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(71) Applicant: MILLENNIUM PHARMACEUTICALS, INC. [US/US]; 640 Memorial Drive, Cambridge, MA 02139

(72) Inventors: LEE, Frank; 6 Norfolk Road, Chestnut Hill, MA 02160 (US). HUSZAR, Dennis; 4 Agawam Road, Acton, MA 01720 (US). GU, Wei; 1080 Beacon Street, Brookline, MA 02146 (US).

(74) Agents: CORUZZI, Laura, A. et al.; Pennie & Edmonds LLP, 1155 Avenue of the Americas, New York, NY 10036 (US). **Published** 

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#### (57) Abstract

The present invention relates to drug screening assays, and diagnostic and therapeutic methods for the treatment of body weight disorders, such as obesity, anorexia and cachexia, utilizing the melanocortin 4-receptor (MC4-R) as the target for intervention. The invention also relates to compounds that modulate the activity or expression of the MC4-R, and the use of such compounds in the treatment of body weight disorders.

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### SCREENING METHODS FOR COMPOUNDS USEFUL IN THE REGULATION OF BODY WEIGHT

#### 1. INTRODUCTION

The present invention relates to drug screening assays, and diagnostic and therapeutic methods for the treatment of body weight disorders, such as obesity, anorexia and cachexia, involving the melanocortin 4-receptor (MC4-R). The invention also relates to compounds that modulate the activity or expression of the MC4-R, and the use of such compounds in the treatment of body weight disorders.

#### 2. BACKGROUND OF THE INVENTION

Melanocortins (a variety of different peptide products resulting from post-translational processing of pro-opiomelanocortin) are known to have a broad array of physiological actions. Aside from their well known effects on adrenal cortical function (e.g., by ACTH, adrenocorticotropic hormone), and on melanocytes (e.g., by 20 α-MSH, melanocyte stimulating hormone), melanocortins have been shown to affect behavior, learning, and memory, control of the cardiovascular system, analgesia, thermoregulation, and the release of other neurohumoral agents including prolactin, luteinizing hormone, and biogenic amines.

25 Peripherally, melanocortins have been identified to have immunomodulatory and neurotrophic properties and to be

involved in events surrounding parturition.

The melanocortins mediate their effects through melanocortin receptors (MC-R) -- a subfamily of G-protein 30 coupled receptors. Other than the MC1-R which was identified as specific for α-MSH, and MC2-R which was identified as specific for ACTH, the melanocortin receptors cloned and identified to date (MC3-R, MC4-R, MC5-R) are thought of as "orphan" receptors -- i.e., the identity of the native ligand 35 for each receptor remains unidentified, and the physiologic function of each receptor type remains unknown.

The agouti protein is a gene product expressed in mice that is known to be involved in determining coat color, but also thought to play a role in obesity when its normal expression pattern is de-regulated and the protein is bubiquitously expressed. The receptor for agouti has not been identified or cloned; however, it has been observed that agouti antagonizes the MSH-induced activation of two melanocortin receptors.

#### 2.1. THE MELANOCORTIN RECEPTORS

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The first two melanocortin receptors cloned were the melanocyte MSH receptor, MC1-R, and the adrenocortical ACTH receptor, MC2-R (Mountjoy et al., 1992, Science 257:1248-1251; Chhajlani & Wikberg, 1992, FEBS Lett. 309:417-420).

- 15 Subsequently, three additional melanocortin receptor genes were cloned which recognize the core heptapeptide sequence (MEHFRWG) of melanocortins. Two of these receptors have been shown to be expressed primarily in the brain, MC3-R (Roselli-Rehfuss et al., 1993, Proc. Natl. Acad. Sci. USA 90:8856-
- 20 8860; Gantz et al., 1993, J. Biol. Chem. 268:8246-8250) and MC4-R (Gantz et al., 1993, J. Biol. Chem. 268:15174-15179; Mountjoy et al., 1994, Mol. Endo. 8:1298-1308). A fifth melanocortin receptor (originally called MC2-R) is expressed in numerous peripheral organs as well as the brain (Chhajlani
- 25 et al., 1993, Biochem. Biophys. Res. Commun. 195:866-873; Gantz et al., 1994, Biochem. Biophs. Res. Commun. 200:1214-1220). The native ligands and functions of these latter three receptors remain unknown.

Because of their "orphan" status as receptors without an identified ligand, and the absence of any known physiological role for these new receptors, investigators have attempted to characterize the receptors in vitro, by their ability to bind and respond (e.g., transduce signal) to a variety of known melanocortins (e.g., see Roselli-Rehfuss, 1993, supra; and

35 Gantz, 1993 <u>supra</u>) or agonists and antagonists deriv d from MSH and ACTH amino acid sequences (<u>e.g.</u>, see Hruby et al., 1995, J. Med. Chem. 38:3454-3461; and Adan et al., 1994, Eur.

J. Pharmacol. 269:331-337). In another approach, the members of the melanocortin receptor family were differentiated on the basis of their pattern of tissue distribution as a means for hypothesizing a function (e.g., See Gantz, 1993, supra; 5 and Mountjoy 1994, supra). For example, expression of MC1-R is localized to melanocytes, MC2-R is localized to adrenal cortical cells, whereas the MC3-R and MC4-R are found primarily in the brain but not in the adrenal cortex or melanocytes; MC4-R is not expressed in the placenta, a tissue 10 that expresses large amounts of MC3-R. Based upon its expression pattern in the hippocampal region of the brain, a role for the MC4-R in learning and memory was proposed (Gantz, 1993, supra) but was noted to be a "pharmacological paradox" in that the MC4-R does not respond well to compounds 15 known to have an effect on retention of learned behaviors. (Mountjoy, 1994, supra). Mountjoy 1994 further suggests that the MC4-R may participate in modulating the flow of visual and sensory information, or coordinate aspects of somatomotor control, and/or may participate in the modulation of 20 autonomic outflow to the heart.

Thus, despite such efforts, the native ligands and function of MC3-R, MC4-R and MC5-R remain elusive.

#### 2.2. THE AGOUTI PROTEIN

- 25 The agouti gene is predicted to encode a secreted protein expressed in hair follicles and the epidermis, the expression of which correlates with the synthesis of the yellow pigment associated with the agouti phenotype (Miller et al., 1993, Gene & Development 7:454-467). Certain
- 30 dominant mutations of the agouti gene result in de-regulated, ubiquitous expression of the agouti protein in mice, demonstrating pleiotropic effects that include obesity and increased tumor susceptibility. (Miller et al., 1993, supra; Michaud et al., 1993, Genes & Development 7:1203-1213).
- 35 Ectopic expression of the normal, wild-type, agouti protein in transgenic mice results in obesity, diabetes, and the yellow coat color commonly observed in spontaneous obese

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mutants (Klebig, et al., 1995, Proc. Natl. Acad. Sci. USA 92:4728-4732). For reviews, see Jackson, 1993, Nature 362:587-588; Conklin & Bourne, 1993, Nature 364:110; Siracusa 1994, TIG 10:423-428; Yen et al., 1994, FASEB J. 8:479-488; 5 Ezzell, 1994, J. NIH Res. 6:31-33; and Manne et al., 1995, Proc. Sci. USA 92:4721-4724.

No receptor for agouti has been identified. Agouti has been reported to be a competitive antagonist of aMSH binding to the MC1-R and MC4-R in vitro (Lu et al., 1994, Nature 10 371:799-802), and the authors speculated that ectopic expression of agouti may lead to obesity by antagonism of melanocortin receptors expressed outside the hair follicle. In this regard, a number of theories have been proposed to account for the induction of obesity by ectopic expression of 15 agouti. For example, agouti expression in skeletal muscle may result in insulin resistance, hyperinsulinemia and obesity via elevation of Ca2+levels; alternatively ectopic agouti expression in adipocytes may depress lipolysis; conversely direct effects of agouti on pancreatic  $\beta$  islet 20 cells may result in hyperinsulinemia and obesity; yet another possibility is that agouti expression in the brain may result in obesity due to a primary effect on areas of the brain controlling weight regulation and insulin production (see Klebig 1995, supra).

In sum, the mechanism of agouti-induced obesity in mice is unknown, and the relevance, if any, of this phenomenon to human obese phenotypes has not been established.

#### 3. SUMMARY OF THE INVENTION

The present invention relates to drug screening assays to identify compounds for the treatment of body weight disorders, such as obesity, anorexia and cachexia by using MC4-R as a target. The invention also relates to compounds that modulate body weight via the MC4-R. The present invention also relates to the treatment of body weight disorders by targeting the MC4-R.

The invention is based, in part, on the discovery of a specific role for MC4-R in body weight regulation. As d monstrated herein, mice completely lacking MC4-R develop a maturity onset obesity syndrome associated with hyperphagia, 5 hyperinsulinemia, and hyperglycemia. In particular, knock-out mice in which the gene encoding MC4-R is defective exhibit significant weight gain compared to either MC4-R heterozygous or wild-type female littermates. The invention is also based, in part, on the discovery that the agouti protein, 10 known to be involved in an obese phenotype when ectopically expressed in mice, binds to the MC4-R.

The invention is further based in part, on the discovery that mutations in the MC4-R have been found to exist in extreme obese human patients. A comparison of the signaling 15 response of the wild type and mutant receptors indicates impaired signaling of the mutant receptor as measured by cAMP induction in the presence of various agonists.

The invention relates to assays designed to screen for compounds or compositions that modulate MC4-R activity, i.e., 20 compounds or compositions that act as agonists or antagonists of MC4-R, and thereby modulate weight control. To this end, cell-based assays or non-cell based assays can be used to detect compounds that interact with, e.g., bind to, a MC4-R extracellular domain ("ECD"). The cell-based assays have the 25 advantage in that they can be used to identify compounds that affect MC4-R biological activity (i.e., signal transduction), including the identification of compounds that do not interact with a MC4-R ECD, but act on an intracellular component of the signal transduction pathway mediated by MC4-30 R.

The invention also relates to assays designed to screen for compounds or compositions that modulate MC4-r gene expression. For example, cell-based assays, or cell-lysate assays (e.g., in vitro transcription or translation assays) can be used to screen for compounds or compositions that modulate MC4-r transcription (e.g., compounds that modulate expression, production or activity of transcription factors

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involved in MC4-r gene expression; polynucleotides that form triple helical structures with an MC4-r regulatory region and inhibit transcription of the MC4-r gene, etc.). Alternatively, cell-based assays or cell-lysate assays can be used to screen for compounds or compositions that modulate translation of MC4-R transcripts (e.g., antisense and ribozyme molecules).

In yet another embodiment, the cell-based assays or cell-lysate assays can be used to test polynucleotide

10 constructs designed to modify the expression of the MC4-r gene in vivo. Such constructs include polynucleotide constructs designed for gene therapy; e.g., expression constructs or gene replacement constructs that place the MC4-r gene under the control of a strong promoter system, an inducible promoter system or a constitutive promoter system.

The invention also encompasses agonists and antagonists of MC4-R, including small molecules, large molecules, and antibodies, as well as nucleotide sequences that can be used to inhibit MC4-r gene expression (e.g., antisense and ribozyme molecules), and gene or regulatory sequence replacement constructs designed to enhance MC4-r gene expression (e.g., expression constructs that place the MC4-r gene under the control of a strong promoter system). Such compounds may be used to treat body weight disorders.

In addition, this invention presents methods for the diagnostic evaluation and prognosis of body weight disorders, including obesity, cachexia and anorexia, and for the identification of subjects having a predisposition to such conditions. For example, nucleic acid molecules encoding

MC4-R can be used as diagnostic hybridization probes or as primers for diagnostic PCR analysis for the identification of MC4-R gene mutations, allelic variations and regulatory defects in the MC4-R gene based in part on the identification of MC4-R mutants in human obese patients.

The invention also encompasses the use of such compounds and compositions, including gene therapy approaches, that

modulate MC4-R activity or MC4-r gene expression to treat body weight disorders.

#### 3.1. **DEFINITIONS**

5 The following terms as used herein shall have the meaning indicated.

MC4-r nucleotides or coding sequences: means DNA sequences encoding MC4-R mRNA transcripts, MC4-R protein, polypeptide or peptide fragments of MC4-R protein, or MC4-R fusion proteins. MC4-r nucleotide sequences encompass DNA, including genomic DNA (e.g. the MC4-r gene) or cDNA.

MC4-R means MC4-r gene products, e.g., transcripts and the MC4 receptor protein. Polypeptides or peptide fragments of the MC4-R protein are referred to as MC4-R polypeptides or 15 MC4-R peptides. Fusions of MC4-R, or MC4-R polypeptides, or peptide fragments to an unrelated protein are referred to herein as MC4-R fusion proteins. A functional MC4-R refers to a protein which binds melanocortin peptides in vivo or in vitro.

20 ECD: means "extracellular domain".

TM: means "transmembrane domain".

CD: means "cytoplasmic domain".

#### 4. DESCRIPTION OF THE FIGURES

- 25 FIG. 1. Deduced amino acid sequences of the melanocortin receptors. The serpentine structure of the melanocortin receptors predicts that the hydrophilic domains located between the TM domains are arranged alternately outside and within the cell to form the ECD (amino acid residues 1-74,
- 30 137-155, 219-231 and 305-316 in FIG. 1) and the CD (amino acid residues 102-112, 178-197, 251-280 and 339-end in FIG. 1) of the receptor. The predicted transmembrane domains are denoted by overbars and Roman numerals, and the four
- extracellular domains (ECD1, ECD2, ECD3 and ECD4) and four
- 35 cytoplasmic domains (CD1, CD2, CD3 and CD4) are indicated.

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- FIG. 2. Schematic diagram of the construction of the MC4-R targeting vector. FIG. 2A. Partial restriction map of the MC4-R locus. FIG. 2B. The MC4 KO 5' construct, containing genomic sequences from 3' of the MC4-R gene in the vector 5 pJN2. FIG. 2C. The MC4-R KO 5'3' construct in which genomic sequences from 5' of the MC4-R gene have been inserted into the MC4 KO 5' construct. FIG. 2D. The MC4-R KO 5'3' neo construct in which a neo expression cassette has been inserted between the 5' and 3' flanking sequences of the MC4-10 R gene. The dotted line represents the pJN2 vector. The open box represents the PGK-neo expression cassette, the hatched box represents the MC4-R gene and the arrows indicate the direction of transcription.
- Schematic diagram of the gene targeting strategy for inactivation of the MC4-R. FIG. 3A. Diagram of the MC4-R The hatched box represents MC4-R coding sequences, the solid box indicates the location of the SacI-SphI probe used for identifying homologous recombinants. The arrow 20 indicates the direction of transcription of the MC4-R gene. FIG. 3B. Diagram of the MC4-R targeting construct. dashed line represents pJN2 plasmid sequences and the arrow indicates the direction of neo transcription. FIG. 3C. Diagram of the MC4-R locus following homologous recombination 25 with the targeting vector. FIG. 3D. Predicted restriction fragment lengths for the wild type and mutated MC4-R loci digested with the indicated enzymes and probed with the SacI-SphI probe. FIG. 3E. Autoradiogram of a Southern blot analysis of tail DNA from F2 progeny. Genomic DNA was 30 digested with ApaI or NcoI, as indicated and hybridized with the radiolabeled probe shown in (A), then stripped and rehybridized with a radiolableed probe consisting of the human MC4-R coding sequence. +/+, +/-, and -/- denote DNA
- human MC4-R coding sequence. +/+, +/-, and -/- denote DNA from wild-type, heterozygous, and homozygous F2 littermates, 35 respectively.

FIG. 4 Weight gain of MC4-R deficient mice and control littermates. Each lin represents the weight gain of an individual mouse. FIG. 4A. Weight gain of female homozygous (-/-) mutant mice (closed squares) and wild type (+/+) F2 5 controls (open circles). The weights of 9 homozygous and 12 control mice were taken at the times indicated. FIG. 4B. Weight gain of female heterozygous (+/-) mutant mice (x) and wild type (+/+) F2 controls (open circles). The weights of 18 heterozygous and 12 control mice were taken at the times 10 indicated. FIG. 4C. Weight gain of male homozygous (-/-) mutant mice (closed squares) and wild-type (+/+) F2 controls (open circles). The weights of 9 homozygous and 17 control mice were taken at the times indicated. FIG. 4D. Weight gain of male heterozygous (+/-) mutant mice (x) and wild-type 15 (+/+) F2 controls (open circles). The weights of 18 heterozygous and 17 control mice were taken at the times indicated.

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- FIG. 5 Sequence of the human MC4-R. Transmembrane domains 20 are underlined. Amino acid differences in the rat MC4-R are indicated underneath the human sequence.
- FIG. 6 Increased linear growth of MC4-R deficient mice. The body length of female (open bars) and male (cross-hatched 25 bars) was measured at approximately 19 weeks of age (between 132-138 days). The bars indicate the mean length of 12 wild type (+/+), and 9 homozygous mutant (-/-) female F2 mice, and 15 wild type, 20 heterozygous, and 9 homozygous mutant male F2 mice. Error bars represent the standard error of the 30 mean, and the asterisks denote significant difference (p<0.02 by two tailed Student t test) compared to the wild type value within a similar sex.
- FIG. 7 Mice lacking the MC4-R are hyperphagic. The food 35 intake of female mice housed in pairs was measured every weekday over a two week period. The open bars represent the mean of 8 measurements on one cage each of two Ay and two

control C57BL/6 mice. The hatched bars represent the mean of 8 measurements on each of two cages of two homozygous mutant mice (-/-) and two F2 wild type controls (+/+). Error bars represent the standard error of the mean, and the asterisks denote significant difference (p<0.01 by two tailed Student t test) of either Ay compared to C57BL/6 or MC4-R (-/-) homozygous mutants compared to MC4-R (+/+) wild type F2 mice.

- FIG. 8. Serum glucose, insulin, and leptin levels in mice 10 lacking the MC4-R. Glucose, insulin and leptin were each measured on the same serum samples. Open bars representheterozygotes, and shaded bars represent homozygous mutant mice. Error bars indicate the standard error of the mean. Asterisks denote significant difference (p<0.05 by two-tailed 15 student t test) compared to control within the same sex and age group. For female mice, the n for wild type mice at 4-8 weeks, 10-14 weeks, and 17-23 weeks was 11, 14, and 7, respectively; and for homozygous mutants, 7, 11 and 3, respectively. For male mice, the n for wild-type mice at 4-8 20 weeks, 10-14 weeks and 17-23 weeks was 14, 14, and 6, respectively; and for homozygous mutants, 8, 8, and 9, respectively. FIG. 8A and 8B. Serum glucose levels of female and male mice, respectively. Five  $\mu l$  of serum was analyzed using a glucose oxidase assay. FIG. 8C and 8D. 25 Serum insulin levels of female and male mice, respectively, were assayed by radioimmunoassay using rat insulin as the standard. FIG. 8E and 8F. Serum leptin levels of female and
- 30 FIG. 9. MC4-R gene deletion does not affect basal serum corticosterone. Serum corticosterone levels were measured in three sets of sex matched littermates containing a representative animal of each genotype:+/+ wild-type control, +/- heterozygote, -/-homozygous mutant. Sets are, from left to right, male, female, and male. Males were 15 weeks of age, females were 18 weeks of age. Data indicate the means of measurements performed using two serum samples obtained on

male mice, respectively, were measured by radioimmunoassay.

different days. Measurement on each day was performed in duplicate. Bars indicate standard deviation. Analysis of data by two-way ANOVA indicated no significant difference in corticosterone levels as a function of genotype.

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- FIG. 10. MC4-R gene deletion does not effect brain POMC mRNA levels. FIG. 10A, 10B and 10C. Hematoxylin and eosin stained brain sections from wild type, heterozygous, and homozygous mutant MC4-R deficient mice, respectively. FIG.
- 10 10D, 10E and 10F. Autoradiographs of brain sections from wild type, heterozygous, and homozygous mutant MC4-R deficient mice, respectively, hybridized with a 35S-POMC antisense cRNA probe.
- 15 FIG. 11A-11B. Sequence of mutant MC4-R. Mutation is Ile137Thr (T to C) mutation.
  - FIG. 12A-12B. Sequence of mutant MC4-R. Mutation is Val102Ile (G to A) mutation.

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- FIG. 13A-13B. Sequence of mutant MC4-R. Mutation is Thr112Met (C to T) mutation.
- FIG. 14. Impaired Signaling of the I137T mutant receptor.
- 25 The signaling response of the wild type (wt) and mutant (mt) receptor to five endogenous melanocortins,  $\alpha$ -MSH (alpha),  $\beta$ -MSH (beta),  $\gamma$ 1-MSH 9 (gamma1),  $\gamma$ 2-MSH (gamma2) and ACTH was compared.

#### 5. <u>DETAILED DESCRIPTION OF THE INVENTION</u>

The invention described in the subsections below encompasses screening methods (e.g., assays) for the identification of compounds which affect weight modulation. The invention also encompasses agonists and antagonists of

35 MC4-R, including small mol cules, large molecules, and antibodies, as well as nucleotide sequences that can be used to inhibit MC4-r gene expression (e.g., antisense and

ribozyme molecules), and gene or regulatory sequence replacement constructs designed to enhance MC4-r gene expression (e.g., expression constructs that place the MC4-r gene under the control of a strong promoter system). Such 5 compounds may be used to treat body weight disorders.

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In particular, cellular and non-cellular assays are described that can be used to identify compounds that interact with the MC4-R, e.g., modulate the activity of the MC4-R and/or bind to the MC4-R. The cell based assays can be 10 used to identify compounds or compositions that affect the signal-transduction activity of the MC4-R, whether they bind to the MC4-R or act on intracellular factors involved in the MC4-R signal transduction pathway. Such cell-based assays of the invention utilize cells, cell lines, or engineered cells or cell lines that express the MC4-R. The cells can be further engineered to incorporate a reporter molecule linked to the signal transduced by the activated MC4-R to aid in the identification of compounds that modulate MC4-R signalling activity.

20 The invention also encompasses the use of cell-based assays or cell-lysate assays (e.g., in vitro transcription or translation assays) to screen for compounds or compositions that modulate MC4-r gene expression. To this end, constructs containing a reporter sequence linked to a regulatory element 25 of the MC4-r gene can be used in engineered cells, or in cell lysate extracts, to screen for compounds that modulate the expression of the reporter gene product at the level of transcription. For example, such assays could be used to identify compounds that modulate the expression or activity 30 of transcription factors involved in MC4-r gene expression, or to test the activity of triple helix polynucleotides. Alternatively, engineered cells or translation extracts can be used to screen for compounds (including antisense and ribozyme constructs) that modulate the translation of MC4-R 35 mRNA transcripts, and therefore, affect expression of the MC4-R.

The invention also encompasses MC4-R proteins, polypeptides (including soluble MC4-R polypeptides or peptides) and MC4-R fusion proteins for use in non-cell based screening assays, for use in generating antibodies, for 5 diagnostics and therapeutics. The MC4-R is predicted to be a serpentine receptor that traverses the membrane seven times, resulting in four extra-cellular domains (ECDs) and four cellular domains (CDs) (see FIG. 1). Peptides corresponding to each ECD, or a polypeptide composed of two or more of the 10 four ECDs linked together can be engineered as described in Section 5.3.1, infra. Alternatively, such peptides or polypeptides can be fused to a heterologous protein, e.g., a reporter, an Ig Fc region, etc., to yield a fusion protein. Such peptides, polypeptides and fusion proteins can be used 15 in the non-cell based assays for screening compounds that interact with, e.g., modulate the activity of the MC4-R and or bind to the MC4-R.

MC4-R protein products can be used to treat weight disorders such as obesity, anorexia or cachexia. Such MC4-R protein products include but are not limited to soluble derivatives such as peptides or polypeptides corresponding to one or more MC4-R ECDs; truncated MC4-R polypeptides lacking one or more ECD or TM; and MC4-R fusion protein products (especially MC4-R-Ig fusion proteins, i.e., fusions of the MC4-R or a domain of the MC4-R, to an IgFc domain). Alternatively, antibodies to the MC4-R or anti-idiotypic antibodies that mimic the MC4-R (including Fab fragments), antagonists or agonists (including compounds that modulate signal transduction which may act on downstream targets in the MC4-R signal transduction pathway) can be used to treat body weight disorders such as obesity, anorexia or cachexia.

For example, the administration of an effective amount of soluble MC4-R polypeptide, or an MC4-R fusion protein (e.g., MC4-R ECD-IgFc) or an anti-idiotypic antibody (or its 35 Fab) that mimics the MC4-R ECD would interact with and thereby "mop up" or "neutralize" endogenous MC4-R ligand, and prevent or reduce binding and receptor activation, leading to

weight gain. In yet another approach, nucleotide constructs encoding such MC4-R products can be used to genetically engineer host cells to express such MC4-R products in vivo; these genetically engineered cells can function as bioreactors in the body delivering a continuous supply of the MC4-R, MC4-R peptide, soluble MC4-R polypeptide, or MC4-R fusion protein that will "mop up" or neutralize MC4-R ligand.

"Gene therapy" approaches for the modulation of MC4-R expression and/or activity in the treatment of body weight 10 disorders are within the scope of the invention. For example, nucleotide constructs encoding functional MC4-Rs, mutant MC4-Rs, as well as antisense and ribozyme molecules can be used to modulate MC4-r expression.

The invention also encompasses pharmaceutical 15 formulations and methods for treating body weight disorders.

### 5.1. THE ROLE OF MC4-R IN THE REGULATION OF BODY WEIGHT

The specific role of the MC4-R protein in vivo was
investigated by engineering MC4-R "knock out" mice in which
most of the endogenous MC4-R gene coding sequence was
deleted, thereby creating mice which are unable to produce
functional MC4-R protein. Unlike MC-R agonist/antagonist
studies which are complicated because each of the MC
receptors, rather than just MC4-R, can be affected, this
specific elimination of only MC4-R function allowed an
evaluation of the biological function of MC4-R.

In order to produce the MC4-R knock out mice, human MC4-r gene sequences were utilized to isolate and clone the murine MC4-r gene. A murine MC4-r targeting construct was then generated which was designed to delete the majority of the murine MC4-r coding sequence upon homologous recombination with the endogenous murine MC4-r gene.

Embryonic stem (ES) cells containing the disrupted MC4-r gene were produced, isolated and microinjected into murine blastocysts to yield mice chimeric for cells containing a disrupted MC4-r gene. Offspring of the chimeric mice

resulting from germlin transmission of the ES genome were obtained and animals heterozygous for the disrupted MC4-R were identified.

In order to assess the role of MC4-R in vivo, the

5 animals heterozygous for the MC4-r disrupted gene were bred
together, producing litters containing wild-type mice, mice
heterozygous for the MC4-r mutation and mice homozygous for
the MC4-R mutation. Inactivation of the MC4-R by gene
targeting results in mice that develop a maturity onset

10 obesity syndrome associated with hyperphagia,
hyperinsulinemia, and hyperglycemia.

The weight gain of the animals was monitored regularly. Homozygous null MC4-R mutants showed an increase in weight compared to mice heterozygous for MC4-R deletion and wild 15 type mice as early as 25 days of age. By approximately 5 weeks of age, most of the homozygous mutants, both males and females, were heavier than their wild type siblings of the same sex, and by 7 weeks of age all of the null mutants were heavier than the controls (FIG. 4A and 4C). By 15 weeks of 20 age, homozygous mutant females were on average twice as heavy as their wild-type siblings, while homozygous mutant males were approximately 50% heavier than wild-type controls. Mice heterozygous for MC4-R deletion showed a weight gain intermediate to that seen in wild-type and homozygous mutant 25 siblings (FIG. 4B and 4D), demonstrating a gene dosage effect of MC4-R ablation on body weight regulation.

In addition, as demonstrated in FIG. 6, MC4-R deficient mice are significantly longer than wild-type controls. The mean length of homozygous mutant females is increased

30 approximately 11% relative to wild-type F2 mice, and heterozygous females are approximately 7% longer than controls. Male homozygotes and heterozygotes are approximately 8% and 2.5% longer than controls, respectively. Absence of the MC4-R also resulted in a significant increase 35 (46%) in food consumption over wild-type F2 controls.

Blood was collected from MC4-R deficient mice and assayed for serum levels of glucose, insulin and leptin

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concentrations. Serum glucose levels were essentially unchanged in females heterozygous or homozygous for MC4-R deletion, but both heterozygous and homozygous males were hyperglycemic (FIG. 8A and 8B). Both male and female mice 5 were also found to be hyperinsulinemic (FIG. 8C and 8D). Heterozygous mutants were hyperinsulinemic, although less so than homozygous mutants. In addition to glucose and insulin, serum leptin levels were altered in MC4-R deficient mice (FIG. 8R and 8F). Heterozygous mice, for the most part, 10 showed leptin levels intermediate between that observed for wild-type mice and homozygous mutants.

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The knock out experiments described herein represent definitive evidence of the role of MC4-R in weight regulation. The experimental design does <u>not</u> rely on the 15 relationship, if any, of the agouti ligand for the characterization of the functional role of the MC4-R.

In addition, the role of MC4-R in weight regulation is demonstrated by the discovery of mutant forms of MC4-R varients in obese human patients. A comparison of the 20 signaling response of the wild type and mutant receptors indicate impaired signaling of the mutant receptor as measured by cAMP induction in the presence of various agonists. Compared with the wildtype receptor, the mutant has much lower maximum activation, i.e., lower maximum cAMP 25 level achieved; and it generally has higher EC50, i.e., higher agonist concentration required to reach half maximum activation. The mutant receptor is only marginally active in the presence of very high agonist concentration that may not be reached under physiological conditions in vivo.

### 5.2. SCREENING ASSAYS FOR DRUGS USEFUL IN REGULATION OF BODY WEIGHT

At least three different assay systems, described in the subsections below, can be designed and used to identify compounds or compositions that modulate MC4-R activity or MC4-r gene expression, and therefore, modulate weight control.

The systems described below may be formulated into kits. To this end, the MC4-R or cells expressing the MC4-R can be packaged in a variety of containers, e.g., vials, tubes, microtitre well plates, bottles, and the like. Other 5 reagents can be included in separate containers and provided with the kit; e.g., positive controls samples, negative control samples, melanocortin peptides (including but not limited to aMSH and ACTH derivatives), buffers, cell culture media, etc.

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#### 5.2.1. CELL-BASED ASSAYS

In accordance with the invention, a cell-based assay system can be used to screen for compounds that modulate the activity of the MC4-R and thereby, modulate body weight. 15 this end, cells that endogenously express MC4-R can be used to screen for compounds. Alternatively, cell lines, such as 293 cells, COS cells, CHO cells, fibroblasts, and the like, genetically engineered to express the MC4-R can be used for screening purposes. Preferably, host cells genetically 20 engineered to express a functional receptor that responds to activation by melanocortin peptides can be used as an endpoint in the assay; e.g., as measured by a chemical, physiological, biological, or phenotypic change, induction of a host cell gene or a reporter gene, change in cAMP levels, 25 adenylyl cyclase activity, host cell G protein activity, extracellular acidification rate, host cell kinase activity, proliferation, differentiation, etc.

In addition, cell-based assay systems can be used to screen for compounds that modulate the activity of mutant 30 MC4-R and thereby, modulate body weight. For example, compounds may be identified which increase the activity of mutant MC4-R thereby alleviating the symptoms of body weight disorders arising from mutant MC4-R. Cell lines, such as 293 cells, COS cells, CHO cells, fibroblasts and the like may be 35 genetically engineered to expr ss mutant receptor. Alternatively, cells that endogenously express mutant MC4 receptor can be used to screen for compounds.

To be useful in screening assays, the host cells expressing functional MC4-R should give a significant response to MC4-R ligand, preferably greater than 5-fold induction over background. Host cells should preferably 5 possess a number of characteristics, depending on the readout, to maximize the inductive response by melanocortin peptides, for example, for detecting a strong induction of a CRE reporter gene: (a) a low natural level of cAMP, (b) G proteins capable of interacting with the MC4-R, (c) a high 10 level of adenylyl cyclase, (d) a high level of protein kinase A, (e) a low level of phosphodiesterases, and (f) a high level of cAMP response element binding protein would be advantageous. To increase response to melanocortin peptide, host cells could be engineered to express a greater amount of 15 favorable factors or a lesser amount of unfavorable factors. In addition, alternative pathways for induction of the CRE reporter could be eliminated to reduce basal levels.

In utilizing such cell systems, the cells expressing the melanocortin receptor are exposed to a test compound or to 20 vehicle controls (e.g., placebos). After exposure, the cells can be assayed to measure the expression and/or activity of components of the signal transduction pathway of the melanocortin receptor, or the activity of the signal transduction pathway itself can be assayed. For example, 25 after exposure, cell lysates can be assayed for induction of cAMP. The ability of a test compound to increase levels of cAMP, above those levels seen with cells treated with a vehicle control, indicates that the test compound induces signal transduction mediated by the melanocortin receptor 30 expressed by the host cell.

To determine intracellular cAMP concentrations, a scintillation proximity assay (SPA) may be utilized (SPA kit is provided by Amersham Life Sciences, Illinois). The assay utilizes 125I label cAMP, an anti-cAMP antibody, and a scintillant-incorporated microsphere coated with a secondary antibody. When brought into close proximity to the microsphere through the labeled cAMP-antibody complex, 125I

will excite the scintillant to emit light. Unlabeled cAMP extracted from cells competes with the <sup>125</sup>I-labeled cAMP for binding to the antibody and thereby diminishes scintillation. The assay may be performed in 96-well plates to enable high-5 throughput screening and 96 well-based scintillation counting instruments such as those manufactured by Wallac or PAckard may be used for readout.

In screening for compounds that may act as antagonists of MC4-R, it is necessary to include ligands that activate 10 the MC4-R, e.g.,  $\alpha$ -MSH,  $\beta$ -MSH or ACTH, to test for inhibition of signal transduction by the test compound as compared to vehicle controls.

In a specific embodiment of the invention, constructs containing the cAMP responsive element linked to any of a 15 variety of different reporter genes may be introduced into cells expressing the melanocortin receptor. Such reporter genes may include but is not limited to chloramphenicol acetyltransferase (CAT), luciferase, GUS, growth hormone, or placental alkaline phosphatase (SEAP). Following exposure of 20 the cells to the test compound, the level of reporter gene expression may be quantitated to determine the test compound's ability to regulate receptor activity. Alkaline phosphatase assays are particularly useful in the practice of the invention as the enzyme is secreted from the cell.

25 Therefore, tissue culture supernatant may be assayed for secreted alkaline phosphatase. In addition, alkaline phosphatase activity may be measured by calorimetric, bioluminescent or chemilumenscent assays such as those described in Bronstein, I. et al. (1994, Biotechniques 17:

30 172-177). Such assays provide a simple, sensitive easily automatable detection system for pharmaceutical screening.

When it is desired to discriminate between the melanocortin receptors and to identify compounds that selectively agonize or antagonize the MC4-R, the assays described above should be conducted using a panel of host cells, each genetically engineered to express one of the melanocortin receptors (MC1-R through MC5-R). Expression of

the human melanocortin receptors is preferred for drug discovery purposes. To this end, host cells can be genetically engineered to express any of the amino acid sequences shown for melanocortin receptors 1 through 5 in The cloning and characterization of each receptor has been described: MC1-R and MC2-R (Mountjoy., 1992, Science 257: 1248-1251; Chhajlani & Wikberg, 1992 FEBS Lett. 309: 417-420); MC3-R (Roselli-Rehfuss et al., 1993, Proc. Natl. Acad. Sci., USA 90: 8856-8860; Gantz et al., 1993, J. 10 Biol. Chem. 268: 8246-8250); MC4-R (Gantz et al., 1993, J. Biol. Chem. 268: 15174-15179; Mountjoy et al., 1994, Mol. Endo. 8: 1298-1308); and MC5-R (Chhajlani et al., 1993, Biochem. Biophys. Res. Commun. 195: 866-873; Gantz et al., 1994, Biochem. Biophys. Res. Commun. 200; 1214-1220), each of 15 which is incorporated by reference herein in its entirety. Thus, each of the foregoing sequences can be utilized to engineer a cell or cell line that expresses one of the melanocortin receptors for use in screening assays described herein. To identify compounds that specifically or 20 selectively regulate MC4-R activity, the activation, or inhibition of MC4-R activation is compared to the effect of the test compound on the other melanocortin receptors. In a specific embodiment, MC1-R through MC5-R cDNAs are expressed in 293 cells under the transcriptional control of 25 the CMV promoter. Stable cell lines are established. transfected human MC2-R (ACTH-R) did not express very well in 293 cells, the human adrenocortical carcinoma cell line H295 (ATCC No. CRL-2128), which expresses endogenous ACTH-R, may be used in screening assays in addition to a stable cell line 30 that expresses transfected ACTH-R. In the first round of screening, the MC4-R expressing cell line is used to identify candidate compounds that activated the MC4-R. identified, those candidate compounds can be tested to determine whether they selectively activate the MC4-R. The 35 activation of the melanocortin receptors may be assayed

using, for example, the SPA assay described above.

Alternatively, if the host cells express more than one melanocortin peptide receptor, the background signal produced by these receptors in response to melanocortin peptides must be "subtracted" from the signal (see Gantz et al., supra).

- 5 The background response produced by these non-MC4-R melanocortin receptors can be determined by a number of methods, including elimination of MC4-R activity by antisense, antibody or antagonist. In this regard, it should be noted that wild type CHO cells demonstrate a small
- 10 endogenous response to melanocortin peptides which must be subtracted from background. Alternatively, activity contributed from other melanocortin receptors could be eliminated by activating host cells with a MC4-R-specific ligand, or including specific inhibitors of the other 15 melanocortin receptors.

#### 5.2.2. NON-CELL BASED ASSAYS

In addition to cell based assays, non-cell based assay systems may be used to identify compounds that interact with, 20 e.g., bind to MC4-R. Such compounds may act as antagonists or agonists of MC4-R activity and may be used in the treatment of body weight disorders.

Isolated membranes may be used to identify compounds that interact with MC4-R. For example, in a typical 25 experiment using isolated membranes, 293 cells may be genetically engineered to express the MC4-R. Membranes can be harvested by standard techniques and used in an <u>in vitro</u> binding assay. <sup>125</sup>I-labelled ligand (e.g., <sup>125</sup>I-labelled α-MSH, β-MSH, or ACTH) is bound to the membranes and assayed for 30 specific activity; specific binding is determined by comparison with binding assays performed in the presence of excess unlabelled ligand.

To identify MC4-R ligands, membranes are incubated with labelled ligand in the presence or absence of test compound.

35 Compounds that bind to the receptor and compete with labelled ligand for binding to the membranes reduced the signal compared to the vehicle control samples.

Alternatively, soluble MC4-R may be recombinantly expressed and utilized in non-cell based assays to identify compounds that bind to MC4-R. The recombinantly expressed MC4-R polypeptides or fusion proteins containing one or more 5 of the ECDs of MC4-R prepared as described in Section 5.3.1, infra, can be used in the non-cell based screening assays. Alternatively, peptides corresponding to one or more of the CDs of MC4-R, or fusion proteins containing one or more of the CDs of MC4-R can be used in non-cell based assay systems 10 to identify compounds that bind to the cytoplasmic portion of the MC4-R; such compounds may be useful to modulate the signal transduction pathway of the MC4-R. In non-cell based assays the recombinantly expressed MC4-R is attached to a solid substrate such as a test tube, microtitre well or a 15 column, by means well known to those in the art (see Ausubel et al., supra). The test compounds are then assayed for their ability to bind to the MC4-R.

In one aspect of the invention the screens may be designed to identify compounds that antagonize the 20 interaction between MC4-R and MC4-R ligands such as  $\alpha$ -MSH,  $\beta$ -MSH and ACTH. In such screens, the MC4-R ligands are labelled and test compounds can be assayed for their ability to antagonize the binding of labelled ligand to MC4-R.

## 5.2.3. ASSAYS FOR COMPOUNDS OR COMPOSITIONS THAT MODULATE EXPRESSION OF THE MC4-R

<u>In vitro</u> cell based assays may be designed to screen for compounds that regulate MC4-R expression at either the transcriptional or translational level.

In one embodiment, DNA encoding a reporter molecule can be linked to a regulatory element of the MC4-r gene and used in appropriate intact cells, cell extracts or lysates to identify compounds that modulate MC4-r gene expression.

Appropriate cells or cell extracts are prepared from any cell type that normally expresses the MC4-r gene, thereby ensuring that the cell extracts contain the transcription factors required for in vitro or in vivo transcription. The screen

can be used to identify compounds that modulate the expression of the reporter construct. In such screens, the level of reporter gene expression is determined in the presence of the test compound and compared to the level of expression in the absence of the test compound.

To identify compounds that regulate MC4-R translation, cells or <u>in vitro</u> cell lysates containing MC4-R transcripts may be tested for modulation of MC4-R mRNA translation. To assay for inhibitors of MC4-R translation, test compounds are 10 assayed for their ability to modulate the translation of MC4-R mRNA in <u>in vitro</u> translation extracts.

Compounds that decrease the level of MC4-R expression, either at the transcriptional or translational level, may be useful for treatment of body weight disorders such as

15 anorexia and cachexia. In contrast, those compounds that increase the expression of MC4-R may be useful for treatment of disorders such as obesity.

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### 5.2.4. COMPOUNDS THAT CAN BE SCREENED IN ACCORDANCE WITH THE INVENTION

The assays described above can identify compounds which affect MC4-R activity. For example, compounds that affect MC4-R activity include but are not limited to compounds that bind to the MC4-R, inhibit binding of the natural ligand, and 25 either activate signal transduction (agonists) or block activation (antagonists), and compounds that bind to the. natural ligand of the MC4-R and neutralize ligand activity. Compounds that affect MC4-r gene activity (by affecting MC4-r gene expression, including molecules, e.g., proteins or small 30 organic molecules, that affect transcription or interfere with splicing events so that expression of the full length or the truncated form of the MC4-R can be modulated) can also be identified on the screens of the invention. However, it should be noted that the assays described can also identify 35 compounds that modulate MC4-R signal transduction (e.g., compounds which affect downstream signalling events, such as inhibitors or enhancers of G protein activities which

participate in transducing the signal activated by ligand binding to the MC4-R). The identification and use of such compounds which affect signalling events downstream of MC4-R and thus modulate effects of MC4-R on the development of body weight disorders are within the scope of the invention. In some instances, G protein-coupled receptors response has been observed to subside, or become desensitized with prolonged exposure to ligand. In an embodiment of the invention assays may be utilized to identify compounds that block the

10 desensitization of the MC4-receptor, such compounds may be used to sustain the activity of the MC4-receptor, such compounds may be used to sustain the activity of the MC4-receptor. Such compounds can be used as part of a therapeutic method for the treatment of body weight

The compounds which may be screened in accordance with the invention include, but are not limited to peptides, antibodies and fragments thereof, and other organic compounds (e.g., peptidomimetics) that bind to the ECD of the MC4-R and either mimic the activity triggered by the natural ligand (i.e., agonists) or inhibit the activity triggered by the natural ligand (i.e., antagonists); as well as peptides, antibodies or fragments thereof, and other organic compounds that include the ECD of the MC4-R (or a portion thereof) and bind to and "neutralize" natural ligand.

Compounds may include, but are not limited to, peptides such as, for example, soluble peptides, including but not limited to members of random peptide libraries; (see, e.g., Lam, K.S. et al., 1991, Nature 354:82-84; Houghten, R. et al., 1991, Nature 354:84-86), and combinatorial chemistry-derived molecular library made of D- and/or L- configuration amino acids, phosphopeptides (including, but not limited to, members of random or partially degenerate, directed phosphopeptide libraries; see, e.g., Songyang, Z. et al., 1993, Cell 72:767-778), antibodies (including, but not limited to, polyclonal, monoclonal, humanized, antiidiotypic, chimeric or single chain antibodies, and FAb,

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F(ab')<sub>2</sub> and FAb expression library fragments, and epitopebinding fragments thereof), and small organic or inorganic molecules.

Other compounds which can be screened in accordance with 5 the invention include but are not limited to small organic molecules that are able to cross the blood-brain barrier, gain entry into an appropriate cell and affect the expression of the MC4-R gene or some other gene involved in the MC4-R signal transduction pathway (e.g., by interacting with the 10 regulatory region or transcription factors involved in gene expression); or such compounds that affect the activity of the MC4-R or the activity of some other intracellular factor involved in the MC4-R signal transduction pathway, such as, for example, the MC4-R associated G protein.

identification of compounds, or the improvement of already identified compounds, that can modulate MC4-R expression or activity. Having identified such a compound or composition, the active sites or regions are identified. Such active sites might typically be ligand binding sites. The active site can be identified using methods known in the art including, for example, from the amino acid sequences of peptides, from the nucleotide sequences of nucleic acids, or from study of complexes of the relevant compound or composition with its natural ligand. In the latter case, chemical or X-ray crystallographic methods can be used to find the active site by finding where on the factor the complexed ligand is found.

Next, the three dimensional geometric structure of the 30 active site is determined. This can be done by known methods, including X-ray crystallography, which can determine a complete molecular structure. On the other hand, solid or liquid phase NMR can be used to determine certain intramolecular distances. Any other experimental method of 35 structure determination can be used to obtain partial or complete geometric structures. The geometric structures may be measured with a complexed ligand, natural or artificial,

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which may increase the accuracy of the active site structure determined.

If an incomplete or insufficiently accurate structure is determined, the methods of computer based numerical modelling 5 can be used to complete the structure or improve its accuracy. Any recognized modelling method may be used, including parameterized models specific to particular biopolymers such as proteins or nucleic acids, molecular dynamics models based on computing molecular motions,

10 statistical mechanics models based on thermal ensembles, or combined models. For most types of models, standard molecular force fields, representing the forces between constituent atoms and groups, are necessary, and can be selected from force fields known in physical chemistry. The incomplete or less accurate experimental structures can serve as constraints on the complete and more accurate structures computed by these modeling methods.

Finally, having determined the structure of the active site, either experimentally, by modeling, or by a 20 combination, candidate modulating compounds can be identified by searching databases containing compounds along with information on their molecular structure. Such a search seeks compounds having structures that match the determined active site structure and that interact with the groups defining the 25 active site. Such a search can be manual, but is preferably computer assisted. These compounds found from this search are potential MC4-R modulating compounds.

Alternatively, these methods can be used to identify improved modulating compounds from an already known

30 modulating compound or ligand. The composition of the known compound can be modified and the structural effects of modification can be determined using the experimental and computer modelling methods described above applied to the new composition. The altered structure is then compared to the active site structure of the compound to determine if an improved fit or interaction results. In this manner systematic variations in composition, such as by varying side

groups, can be quickly evaluated to obtain modified

modulating compounds or ligands of improved specificity or activity.

Further experimental and computer modeling methods

Further experimental and computer modeling methods
5 useful to identify modulating compounds based upon
identification of the active sites of MC4-R, and related
transduction and transcription factors will be apparent to
those of skill in the art.

Examples of molecular modelling systems are the CHARMm

10 and QUANTA programs (Polygen Corporation, Waltham, MA).

CHARMm performs the energy minimization and molecular dynamics functions. QUANTA performs the construction, graphic modelling and analysis of molecular structure.

QUANTA allows interactive construction, modification,

15 visualization, and analysis of the behavior of molecules with each other.

A number of articles review computer modelling of drugs interactive with specific proteins, such as Rotivinen, et al.) 1988, Acta Pharmaceutical Fennica 97:159-166); Ripka 20 (1988 New Scientist 54-57); McKinaly and Rossmann (1989, Annu. Rev. Pharmacol. Toxiciol. 29:111-122); Perry and Davies, OSAR: Quantitative Structure-Activity Relationships in Drug Design pp. 189-193 Alan R. Liss, Inc. 1989; Lewis and Dean (1989, Proc. R. Soc. Lond. 236:125-140 and 141-162); 25 and, with respect to a model receptor for nucleic acid components, Askew, et al. (1989, J. Am. Chem. Soc. 111:1082-1090). Other computer programs that screen and graphically depict chemicals are available from companies such as BioDesign, Inc. (Pasadena, CA.), Allelix, Inc. (Mississauga, 30 Ontario, Canada), and Hypercube, Inc. (Cambridge, Ontario). Although these are primarily designed for application to drugs specific to particular proteins, they can be adapted to design of drugs specific to regions of DNA or RNA, once that

Although described above with reference to design and generation of compounds which could alter binding, one could also screen libraries of known compounds, including natural

region is identified.

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products or synthetic chemicals, and biologically active materials, including proteins, for compounds which are inhibitors or activators.

Compounds identified via assays such as those described 5 herein may be useful, for example, in elaborating the biological function of the MC4-R gene product, and for ameliorating body weight disorders. Assays for testing the efficacy of compounds identified in the cellular screen can be tested in animal model systems for body weight disorders. 10 Such animal models may be used as test substrates for the identification of drugs, pharmaceuticals, therapies and interventions which may be effective in treating such disorders. For example, animal models may be exposed to a compound, suspected of exhibiting an ability to ameliorate 15 body weight disorder symptoms, at a sufficient concentration and for a time sufficient to elicit such an amelioration of body weight disorder symptoms in the exposed animals. response of the animals to the exposure may be monitored by assessing the reversal of disorders associated with body 20 weight disorders such as obesity. With regard to intervention, any treatments which reverse any aspect of body weight disorder-like symptoms should be considered as candidates for human body weight disorder therapeutic intervention. Dosages of test agents may be determined by 25 deriving dose-response curves, as discussed in Section 5.5,

To this end, transgenic animals that express the human MC4-r gene products can be used. Animals of any species, including, but not limited to, mice, rats, rabbits, guinea 30 pigs, pigs, micro-pigs, goats, and non-human primates, e.g., baboons, monkeys, and chimpanzees may be used to generate MC4-R transgenic animals.

below.

Any technique known in the art may be used to introduce the human MC4-r transgene into animals to produce the founder 35 lin s of transgenic animals. Such techniques include, but are not limited to pronuclear microinjection (Hopp, P.C. and Wagner, 1989, U.S. Pat. No. 4,873,191); retrovirus mediated

gene transfer into germ lines (Van der Putten et al., 1985, Proc. Natl. Acad. Sci., USA 82:6148-6152); gene targeting in embryonic stem cells (Thompson et al., 1989, Cell 56:313-321); electroporation of embryos (Lo, 1983, Mol Cell. Biol.

- 5 3:1803-1814); and sperm-mediated gene transfer (Lavitrano et al., 1989, Cell 57:717-723); etc. For a review of such techniques, see Gordon, 1989, Transgenic Animals, Intl. Rev. Cytol. 115:171-229, which is incorporated by reference herein in its entirety.
- The present invention provides for transgenic animals that carry the MC4-r transgene in all their cells, as well as animals which carry the transgene in some, but not all their cells, i.e., mosaic animals. The transgene may be integrated as a single transgene or in concatamers, e.g., head-to-head
- 15 tandems or head-to-tail tandems. The transgene may also be selectively introduced into and activated in a particular cell type by following, for example, the teaching of Lasko et al. (Lasko, M. et al., 1992, Proc. Natl. Acad. Sci. USA 89: 6232-6236). The regulatory sequences required for such a
- 20 cell-type specific activation will depend upon the particular cell type of interest, and will be apparent to those of skill in the art. When it is desired that the MC4-r transgene be integrated into the chromosomal site of the endogenous MC4-r gene, gene targeting is preferred. Briefly, when such a
- 25 technique is to be utilized, vectors containing nucleotide sequences homologous to the endogenous MC4-r gene and/or sequences flanking the gene are designed for the purpose of integrating, via homologous recombination with chromosomal sequences, into and disrupting the function of the endogenous
- 30 MC4-r gene. The transgene may also be selectively expressed in a particular cell type with concomitant inactivation of the endogenous MC4-r gene in only that cell type, by following, for example, the teaching of Gu et al. (Gu, et al., 1994, Science 265: 103-106). The regulatory sequences
- 35 required for such a cell-type specific recombination will depend upon the particular cell type of interest, and will be apparent to those of skill in the art.

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Once founder animals have been generated, standard techniques such as Southern blot analysis or PCR techniques are used to analyze animal tissues to determine whether integration of the transgene has taken place. The level of 5 mRNA expression of the transgene in the tissues of the founder animals may also be assessed using techniques which include but are not limited to Northern blot analysis of tissue samples obtained from the animal, in situ hybridization analysis, and RT-PCR. Samples of MC4-R gene
10 expressing tissue, may also be evaluated immunocytochemically using antibodies specific for the MC4-R transgene product.

#### 5.3. MC4-R PROTEINS, POLYPEPTIDES, AND ANTIBODIES

MC4-R protein, polypeptides and peptide fragments,

15 mutated, truncated or deleted forms of the MC4-R and/or MC4-R
fusion proteins can be prepared for a variety of uses,
including but not limited to the generation of antibodies, as
reagents in diagnostic assays, the identification of other
cellular gene products involved in the regulation of body

20 weight, as reagents in assays for screening for compounds
that can be used in the treatment of body weight disorders,
and as pharmaceutical reagents useful in the treatment of
body weight disorders related to the MC4-R.

#### 25 5.3.1. PRODUCTION OF MC4-R POLYPEPTIDES

The deduced amino acid sequences of the melanocortin receptors, including MC4-R, are shown in FIG. 1, where predicted transmembrane domains are denoted by overbars and Roman numerals, and the four extracellular domains (ECD1, 30 ECD2, ECD3, and ECD4) and the four cytoplasmic domains (CD1, CD2, CD3 and CD4) are indicated. The serpentine structure of the melanocortin receptors predicts that the hydrophilic domains located between the TM domains are arranged alternately outside and within the cell to form the ECD 35 (amino acid residues 1-74, 137-155, 219-231 and 305-316 in FIG. 1) and the CD (amino acid residues 102-112, 178-197, 251-280 and 339-end in FIG. 1) of the receptor. Peptides

corresponding to one or more domains of the MC4-R (e.g., ECDs, TMs or CDs), truncated or deleted MC4-R (e.g., MC4-R in which one or more of the ECDs, TMs and/or CDs is deleted) as well as fusion proteins in which the full length MC4-R, an 5 MC4-R peptide or truncated MC4-R is fused to an unrelated protein are also within the scope of the invention. soluble peptides, proteins, fusion proteins, or antibodies (including anti-idiotypic antibodies) that bind to and "neutralize" circulating natural ligand for the MC4-R, can be 10 used as described in Section 5,5, infra, to effectuate weight gain. To this end, peptides corresponding to individual ECDs of MC4-R, soluble deletion mutants of MC4-R (e.g., ATM mutants), or the entire MC4-R ECD (engineered by linking the four ECDs together as described below) can be fused to 15 another polypeptide (e.g., an IgFc polypeptide). Fusion of the MC4-R or the MC4-R ECD to an IgFc polypeptide should not only increase the stability of the preparation, but will increase the half-life and activity of the MC4-R-Ig fusion protein in vivo. The Fc region of the Ig portion of the 20 fusion protein may be further modified to reduce immunoglobulin effector function.

Such peptides, polypeptides, and fusion proteins can be prepared by recombinant DNA techniques. For example, nucleotide sequences encoding one or more of the four domains 25 of the ECD of the serpentine MC4-R can be synthesized or cloned and ligated together to encode a soluble ECD of the MC4-R. The DNA sequence encoding one or more of the four ECDs (ECD1-4 in FIG. 1) can be ligated together directly or via a linker oligonucleotide that encodes a peptide spacer. 30 Such linkers may encode flexible, glycine-rich amino acid sequences thereby allowing the domains that are strung together to assume a conformation that can bind MC4-R ligands. Alternatively, nucleotide sequences encoding individual domains within the ECD can be used to express MC4-35 R peptides. In addition, mutant MC4-R proteins such as those shown in FIGS. 11-13 can be expressed by recombinant DNA techniques.

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A variety of host-expression vector syst ms may be utilized to express nucleotide sequences encoding the appropriate regions of the MC4-R to produce such polypeptides. Where the resulting peptide or polypeptide is 5 a soluble derivative (e.g., peptides corresponding to the ECDs; truncated or deleted in which the TMs and/or CDs are deleted) the peptide or polypeptide can be recovered from the culture media. Where the polypeptide or protein is not secreted, the MC4-R product can be recovered from the host 10 cell itself.

The host-expression vector systems also encompass engineered host cells that express the MC4-R or functional equivalents in situ, i.e., anchored in the cell membrane. Purification or enrichment of the MC4-R from such expression 15 systems can be accomplished using appropriate detergents and lipid micelles and methods well known to those skilled in the art. However, such engineered host cells themselves may be used in situations where it is important not only to retain the structural and functional characteristics of the MC4-R, 20 but to assess biological activity, e.g., in drug screening assays.

Alternatively, host-expression vector systems may be used to engineer host cells that express mutant MC4-R protein (see, for example FIG. 11-13). Such host cells may be used to assess biological activity, e.g., in drug screening assays.

The host-expression vector systems that may be used for purposes of the invention include but are not limited to microorganisms such as bacteria (e.g., E. coli, B. subtilis)

30 transformed with recombinant bacteriophage DNA, plasmid DNA or cosmid DNA expression vectors containing MC4-R nucleotide sequences; yeast (e.g., Saccharomyces, Pichia) transformed with recombinant yeast expression vectors containing the MC4-R nucleotide sequences; insect cell systems infected with

35 recombinant virus expression vectors (e.g., baculovirus) containing the MC4-R sequences; plant cell systems infect d with recombinant virus expression vectors (e.g., cauliflower

mosaic virus, CaMV; tobacco mosaic virus, TMV) or transformed with recombinant plasmid expression vectors (e.g., Ti plasmid) containing MC4-R nucleotide sequences; or mammalian cell systems (e.g., COS, CHO, BHK, 293, 3T3) harboring recombinant expression constructs containing promoters derived from the genome of mammalian cells (e.g., metallothionein promoter) or from mammalian viruses (e.g., the adenovirus late promoter; the vaccinia virus 7.5K promoter).

In bacterial systems, a number of expression vectors may 10 be advantageously selected depending upon the use intended for the MC4-R gene product being expressed. For example, when a large quantity of such a protein is to be produced, for the generation of pharmaceutical compositions of MC4-R 15 protein or for raising antibodies to the MC4-R protein, for example, vectors which direct the expression of high levels of fusion protein products that are readily purified may be desirable. Such vectors include, but are not limited, to the E. coli expression vector pUR278 (Ruther et al., 1983, EMBO 20 J. 2:1791), in which the MC4-R coding sequence may be ligated individually into the vector in frame with the <a href="lacz">lacz</a> coding region so that a fusion protein is produced; pIN vectors (Inouye & Inouye, 1985, Nucleic Acids Res. 13:3101-3109; Van Heeke & Schuster, 1989, J. Biol. Chem. 264:5503-5509); and 25 the like. pGEX vectors may also be used to express foreign polypeptides as fusion proteins with glutathione S-transferase (GST). In general, such fusion proteins are soluble and can easily be purified from lysed cells by adsorption to glutathione-agarose beads followed by elution 30 in the presence of free glutathione. The PGEX vectors are designed to include thrombin or factor Xa protease cleavage sites so that the cloned target gene product can be released from the GST moiety.

Alternatively, any fusion protein may be readily

35 purified by utilizing an antibody specific for the fusion protein being expressed. For example, a system described by Janknecht et al. allows for the ready purification of non-

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denatured fusion proteins expressed in human cell lines (Janknecht, et al., 1991, Proc. Natl. Acad. Sci. USA 88: 8972-8976). In this system, the gene of interest is subcloned into a vaccinia recombination plasmid such that the 5 gene's open reading frame is translationally fused to an amino-terminal tag consisting of six histidine residues. Extracts from cells infected with recombinant vaccinia virus are loaded onto Ni<sup>2\*</sup> nitriloacetic acid-agarose columns and histidine-tagged proteins are selectively eluted with 10 imidazole-containing buffers.

In an insect system, <u>Autographa californica</u> nuclear polyhedrosis virus (AcNPV) is used as a vector to express foreign genes. The virus grows in <u>Spodoptera frugiperda</u> cells. The MC4-R coding sequence may be cloned individually into non-essential regions (for example the polyhedrin gene) of the virus and placed under control of an AcNPV promoter (for example the polyhedrin promoter). Successful insertion of MC4-R gene coding sequence will result in inactivation of the polyhedrin gene and production of non-occluded recombinant virus (i.e., virus lacking the proteinaceous coat coded for by the polyhedrin gene). The recombinant viruses are then used to infect cells in which the inserted gene is

expressed. (E.g., see Smith et al., 1983, J. Virol. 46: 584;

In mammalian host cells, a number of viral-based expression systems may be utilized. In cases where an adenovirus is used as an expression vector, the MC4-R nucleotide sequence of interest may be ligated to an adenovirus transcription/translation control complex, e.g.,

Smith, U.S. Patent No. 4,215,051).

30 the late promoter and tripartite leader sequence. This chimeric gene may then be inserted in the adenovirus genome by in vitro or in vivo recombination. Insertion in a non-essential region of the viral genome (e.g., region E1 or E3) will result in a recombinant virus that is viable and capable

35 of expressing the MC4-R gene product in infected hosts. (E.g., See Logan & Shenk, 1984, Proc. Natl. Acad. Sci. USA 81:3655-3659). Specific initiation signals may also be

required for efficient translation of inserted MC4-R nucleotide sequences. These signals include the ATG initiation codon and adjacent sequences. In cases where an entire MC4-R gene or cDNA, including its own initiation codon 5 and adjacent sequences, is inserted into the appropriate expression vector, no additional translational control signals may be needed. However, in cases where only a portion of the MC4-R coding sequence is inserted, exogenous translational control signals, including, perhaps, the ATG 10 initiation codon, must be provided. Furthermore, the initiation codon must be in frame with the reading frame of the desired coding sequence to ensure translation of the entire insert. These exogenous translational control signals and initiation codons can be of a variety of origins, both 15 natural and synthetic. The efficiency of expression may be enhanced by the inclusion of appropriate transcription enhancer elements, transcription terminators, etc. (See Bittner et al., 1987, Methods in Enzymol. 153:516-544).

In addition, a host cell strain may be chosen which 20 modulates the expression of the inserted sequences, or modifies and processes the gene product in the specific fashion desired. Such modifications (e.g., glycosylation) and processing (e.g., cleavage) of protein products may be important for the function of the protein. Different host 25 cells have characteristic and specific mechanisms for the post-translational processing and modification of proteins and gene products. Appropriate cell lines or host systems can be chosen to ensure the correct modification and processing of the foreign protein expressed. Accordingly, 30 eukaryotic host cells which possess the cellular machinery for proper processing of the primary transcript, glycosylation, and phosphorylation of the gene product may be used. Such mammalian host cells include, but are not limited to, CHO, VERO, BHK, HeLa, COS, MDCK, 293, 3T3 and WI38 cell 35 lines.

For long-term, high-yield production of recombinant proteins, stable expression is preferred. For example, cell

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lines which stably express the MC4-R sequences described above may be engineered. Rather than using expression vectors which contain viral origins of replication, host cells can be transformed with DNA controlled by appropriate 5 expression control elements (e.g., promoter, enhancer sequences, transcription terminators, polyadenylation sites, etc.), and a selectable marker. Following the introduction of the foreign DNA, engineered cells may be allowed to grow for 1-2 days in an enriched media, and then are switched to a 10 selective media. The selectable marker in the recombinant plasmid confers resistance to the selection and allows cells to stably integrate the plasmid into their chromosomes and grow to form foci which in turn can be cloned and expanded into cell lines. This method may advantageously be used to 15 engineer cell lines which express the MC4-R gene product. Such engineered cell lines may be particularly useful in screening and evaluation of compounds that affect the endogenous activity of the MC4-R gene product.

A number of selection systems may be used, including but 20 not limited to the herpes simplex virus thymidine kinase (Wigler, et al., 1977, Cell 11:223), hypoxanthine-guanine phosphoribosyltransferase (Szybalska & Szybalski, 1962, Proc. Natl. Acad. Sci. USA 48:2026), and adenine phosphoribosyltransferase (Lowy, et al., 1980, Cell 22:817) 25 genes can be employed in tk', hgprt or aprt cells, respectively. Also, antimetabolite resistance can be used as the basis of selection for the following genes: dhfr, which confers resistance to methotrexate (Wigler, et al., 1980, Natl. Acad. Sci. USA 77:3567; O'Hare, et al., 1981, Proc. 30 Natl. Acad. Sci. USA 78:1527); gpt, which confers resistance to mycophenolic acid (Mulligan & Berg, 1981, Proc. Natl. Acad. Sci. USA 78:2072); neo, which confers resistance to the aminoglycoside G-418 (Colberre-Garapin, et al., 1981, J. Mol. Biol. 150:1); and hygro, which confers resistance to 35 hygromycin (Santerre, et al., 1984, Gene 30:147).

## 5.3.2. ANTIBODIES TO MC4-R POLYPEPTIDES

Antibodies that specifically recognize one or more epitopes of MC4-R, or epitopes of conserved variants of MC4-R, or peptide fragments of the MC4-R are also encompassed by the invention. Further, antibodies that specifically

- 5 recognize mutant forms of MC4-R, such as those encoded by the DNA sequences shown in Figures 11-13, are encompassed by the invention. Such antibodies include but are not limited to polyclonal antibodies, monoclonal antibodies (mAbs), humanized or chimeric antibodies, single chain antibodies,
- 10 Fab fragments, F(ab')<sub>2</sub> fragments, fragments produced by a Fab expression library, anti-idiotypic (anti-Id) antibodies, and epitope-binding fragments of any of the above.

The antibodies of the invention may be used, for example, in the detection of the MC4-R in a biological sample 15 and may, therefore, be utilized as part of a diagnostic or prognostic technique whereby patients may be tested for abnormal amounts of MC4-R. Antibodies that specifically recognize mutant forms of MC4-R, such as those described below, in Section 8, may be particularly useful as part of a 20 diagnostic or prognostic technique. Such antibodies may also be utilized in conjunction with, for example, compound screening schemes, as described, above, for the evaluation of the effect of test compounds on expression and/or activity of

25 used in conjunction with the gene therapy techniques described, below, e.g., to evaluate the normal and/or engineered MC4-R-expressing cells prior to their introduction into the patient. Such antibodies may additionally be used as a method for the inhibition of abnormal MC4-R activity.

the MC4-R gene product. Additionally, such antibodies can be

30 Thus, such antibodies may, therefore, be utilized as part of weight disorder treatment methods.

For the production of antibodies, various host animals may be immunized by injection with the MC4-R, an MC4-R peptide (e.g., one corresponding the a functional domain of the receptor, such as ECD, TM or CD), truncated MC4-R polypeptides (MC4-R in which one or more domains, e.g., the TM or CD, has been deleted), functional equivalents of the

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MC4-R or mutants of the MC4-R. Such host animals may include but are not limited to rabbits, mice, hamsters and rats, to name but a few. Various adjuvants may be used to increase the immunological response, depending on the host species;

- 5 including but not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanin, dinitrophenol, and potentially useful human
- 10 adjuvants such as BCG (bacille Calmette-Guerin) and

  Corynebacterium parvum. Polyclonal antibodies are
  heterogeneous populations of antibody molecules derived from
  the sera of the immunized animals.

Monoclonal antibodies, which are homogeneous populations of antibodies to a particular antigen, may be obtained by any technique which provides for the production of antibody molecules by continuous cell lines in culture. These include, but are not limited to, the hybridoma technique of Kohler and Milstein, (1975, Nature 256:495-497; and U.S.

- 20 Patent No. 4,376,110), the human B-cell hybridoma technique (Kosbor et al., 1983, Immunology Today 4:72; Cole et al., 1983, Proc. Natl. Acad. Sci. USA 80:2026-2030), and the EBV-hybridoma technique (Cole et al., 1985, Monoclonal Antibodies And Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). Such
- 25 antibodies may be of any immunoglobulin class including IgG, IgM, IgE, IgA, IgD and any subclass thereof. The hybridoma producing the mAb of this invention may be cultivated in vitro or in vivo. Production of high titers of mAbs in vivo makes this the presently preferred method of production.
- In addition, techniques developed for the production of "chimeric antibodies" (Morrison et al., 1984, Proc. Natl. Acad. Sci., 81:6851-6855; Neuberger et al., 1984, Nature, 312:604-608; Takeda et al., 1985, Nature, 314:452-454) by splicing the genes from a mouse antibody molecule of
- 35 appropriate antigen specificity together with genes from a human antibody molecule of appropriate biological activity can be used. A chimeric antibody is a molecule in which

different portions are derived from different animal species, such as those having a variable region derived from a murine mAb and a human immunoglobulin constant region.

Alternatively, techniqu s described for the production 5 of single chain antibodies (U.S. Patent 4,946,778; Bird, 1988, Science 242:423-426; Huston et al., 1988, Proc. Natl. Acad. Sci. USA 85:5879-5883; and Ward et al., 1989, Nature 334:544-546) can be adapted to produce single chain antibodies against MC4-R gene products. Single chain 10 antibodies are formed by linking the heavy and light chain fragments of the Fv region via an amino acid bridge, resulting in a single chain polypeptide.

Antibody fragments which recognize specific epitopes may be generated by known techniques. For example, such 15 fragments include but are not limited to: the F(ab'), fragments which can be produced by pepsin digestion of the antibody molecule and the Fab fragments which can be generated by reducing the disulfide bridges of the F(ab'), fragments. Alternatively, Fab expression libraries may be 20 constructed (Huse et al., 1989, Science, 246:1275-1281) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity.

Antibodies to the MC4-R can, in turn, be utilized to generate anti-idiotype antibodies that "mimic" the MC4-R,

25 using techniques well known to those skilled in the art.

(See, e.g., Greenspan & Bona, 1993, FASEB J 7(5):437-444; and Nissinoff, 1991, J. Immunol. 147(8):2429-2438). For example antibodies which bind to the MC4-R ECD and competitively inhibit the binding of melanocortins to the MC4-R can be used to generate anti-idiotypes that "mimic" the ECD and, therefore, bind and neutralize melanocortins. Such neutralizing anti-idiotypes or Fab fragments of such anti-idiotypes can be used in therapeutic regimens to neutralize the native ligand and promote weight gain.

Alternatively, antibodies to MC4-R that can act as agonists of MC4-R activity can be generated. Such antibodies will bind to the MC4-R and activate the signal transducing

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activity of the receptor. Such antibodies would be particularly useful for treating weight disorders such as obesity. In addition, antibodies that act as antagonist of MC4-R activity, <u>i.e.</u> inhibit the activation of MC4-R receptor may be used to treat weight disorders such as anorexia or cachexia.

## 5.4. GENE THERAPY APPROACHES TO CONTROLLING MC4-R ACTIVITY AND REGULATING BODY WEIGHT

The expression of MC4-R can be controlled in vivo (e.g. at the transcriptional or translational level) using gene therapy approaches to regulate MC4-R activity and treat body weight disorders. Certain approaches are described below.

#### 5.4.1. GENE REPLACEMENT THERAPY

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with respect to an increase in the level of normal MC4-R gene expression and/or MC4-R gene product activity, MC4-R nucleic acid sequences can be utilized for the treatment of body weight disorders, including obesity. Where the cause of obesity is a defective MC4-R gene, treatment can be administered, for example, in the form of gene replacement therapy. Specifically, one or more copies of a normal MC4-R gene or a portion of the MC4-R gene that directs the production of an MC4-R gene product exhibiting normal function, may be inserted into the appropriate cells within a patient or animal subject, using vectors which include, but are not limited to adenovirus, adeno-associated virus, retrovirus and herpes virus vectors, in addition to other particles that introduce DNA into cells, such as liposomes.

Because the MC4-R gene is expressed in the brain, including the cortex, thalamus, brain stem and spinal cord and hypothalamus, such gene replacement therapy techniques should be capable of delivering MC4-R gene sequences to these cell types within patients. Thus, the techniques for delivery of the MC4-R gene sequences should be designed to readily cross the blood-brain barrier, which are well known to those of skill in the art (see, e.g., PCT application,

publication No. W089/10134, which is incorporated herein by reference in its entirety), or, alternatively, should involve direct administration of such MC4-R gene sequences to the site of the cells in which the MC4-R gene sequences are to be 5 expressed.

Alternatively, targeted homologous recombination can be utilized to correct the defective endogenous MC4-R gene in the appropriate tissue; e.g., brain tissue. In animals, targeted homologous recombination can be used to correct the 10 defect in ES cells in order to generate offspring with a corrected trait.

Additional methods which may be utilized to increase the overall level of MC4-R gene expression and/or MC4-R activity include the introduction of appropriate MC4-R-expressing 15 cells, preferably autologous cells, into a patient at positions and in numbers which are sufficient to ameliorate the symptoms of body weight disorders, including obesity. Such cells may be either recombinant or non-recombinant. Among the cells which can be administered to increase the 20 overall level of MC4-R gene expression in a patient are normal cells, or hypothalamus cells which express the MC4-R gene. The cells can be administered at the anatomical site in the brain, or as part of a tissue graft located at a different site in the body. Such cell-based gene therapy 25 techniques are well known to those skilled in the art, see, e.g., Anderson, et al., U.S. Patent No. 5,399,349; Mulligan & Wilson, U.S. Patent No. 5,460,959.

Finally, compounds, identified in the assays described above, that stimulate or enhance the signal transduced by 30 activated MC4-R, e.g., by activating downstream signalling proteins in the MC4-R cascade and thereby by-passing the defective MC4-R, can be used to achieve weight loss. The formulation and mode of administration will depend upon the physico-chemical properties of the compound. The 35 administration should include known techniques that allow for a crossing of the blood-brain barrier.

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## 5.4.2. <u>INHIBITION OF MC4-R EXPRESSION</u>

In an alternate embodiment, weight gain therapy can be designed to reduce the level of endogenous MC4-R gene expression, e.g., using antisense or ribozyme approaches to 5 inhibit or prevent translation of MC4-R mRNA transcripts; triple helix approaches to inhibit transcription of the MC4-R gene; or targeted homologous recombination to inactivate or "knock out" the MC4-R gene or its endogenous promoter. gene therapy may be utilized for treatment of body weight 10 disorders such as cachexia and anorexia where the inhibition of MC4-R expression is designed to increase body weight. Because the MC4-R gene is expressed in the brain, delivery techniques should be preferably designed to cross the bloodbrain barrier (see PCT WO89/10134, which is incorporated by 15 reference herein in its entirety). Alternatively, the antisense, ribozyme or DNA constructs described herein could ' be administered directly to the site containing the target cells.

Antisense approaches involve the design of 20 oligonucleotides (either DNA or RNA) that are complementary to mRNA. The antisense oligonucleotides will bind to the complementary mRNA transcripts and prevent translation. Absolute complementarily, although preferred, is not required. A sequence "complementary" to a portion of an RNA, 25 as referred to herein, means a sequence having sufficient complementarily to be able to hybridize with the RNA, forming a stable duplex; in the case of double-stranded antisense nucleic acids, a single strand of the duplex DNA may thus be tested, or triplex formation may be assayed. The ability to 30 hybridize will depend on both the degree of complementarily and the length of the antisense nucleic acid. Generally, the longer the hybridizing nucleic acid, the more base mismatches with an RNA it may contain and still form a stable duplex (or triplex, as the case may be). One skilled in the art can 35 ascertain a tolerable degree of mismatch by use of standard procedures to determine the melting point of the hybridized complex.

While antisense nucleotides complementary to the coding region sequence could be used, those complementary to the transcribed untranslated region are most preferred.

Oligonucleotides that are complementary to the 5' end of the message, e.g., the 5' untranslated sequence up to and including the AUG initiation codon, should work most efficiently at inhibiting translation (see FIG. 5). However, sequences complementary to the 3' untranslated sequences of mRNAs have recently shown to be effective at inhibiting translation of mRNAs as well. See generally, Wagner, R., 1994, Nature 372:333-335. Thus, oligonucleotides complementary to either the 5'- or 3'- non- translated, non-coding regions of MC4-R could be used in an antisense

of the mRNA should include the complement of the AUG start codon. Antisense oligonucleotides complementary to mRNA coding regions are less efficient inhibitors of translation but could be used in accordance with the invention. Whether

approach to inhibit translation of endogenous mRNA.

20 designed to hybridize to the 5'-, 3'- or coding region of MC4-R mRNA, antisense nucleic acids should be at least six nucleotides in length, and are preferably oligonucleotides ranging from 6 to about 50 nucleotides in length. In specific aspects the oligonucleotide is at least 10

25 nucleotides, at least 17 nucleotides, at least 25 nucleotides or at least 50 nucleotides.

Regardless of the choice of target sequence, it is preferred that in vitro studies are first performed to quantitate the ability of the antisense oligonucleotide to 30 inhibit gene expression. It is preferred that these studies utilize controls that distinguish between antisense gene inhibition and nonspecific biological effects of oligonucleotides. It is also preferred that these studies compare levels of the target RNA or protein with that of an 35 internal control RNA or protein. Additionally, it is envisioned that results obtain d using the antisense oligonucleotide are compared with those obtained using a

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control oligonucleotide. It is preferred that the control oligonucl otide is of approximately the same length as the test oligonucleotide and that the nucleotide sequence of the oligonucleotide differs from the antisense sequence no more than is nec ssary to prevent specific hybridization to the target sequence.

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The oligonucleotides can be DNA or RNA or chimeric mixtures or derivatives or modified versions thereof, singlestranded or double-stranded. The oligonucleotide can be 10 modified at the base moiety, sugar moiety, or phosphate backbone, for example, to improve stability of the molecule, hybridization, etc. The oligonucleotide may include other appended groups such as peptides (e.g., for targeting host cell receptors in vivo), or agents facilitating transport 15 across the cell membrane (see, e.g., Letsinger et al., 1989, Proc. Natl. Acad. Sci. U.S.A. 86:6553-6556; Lemaitre et al., 1987, Proc. Natl. Acad. Sci. 84:648-652; PCT Publication No. WO88/09810, published December 15, 1988) or the blood-brain barrier (see, e.g., PCT Publication No. WO89/10134, published 20 April 25, 1988), hybridization-triggered cleavage agents. (See, e.g., Krol et al., 1988, BioTechniques 6:958-976) or intercalating agents. (See, e.g., Zon, 1988, Pharm. Res. 5:539-549). To this end, the oligonucleotide may be conjugated to another molecule, e.g., a peptide, 25 hybridization triggered cross-linking agent, transport agent, hybridization-triggered cleavage agent, etc.

The antisense oligonucleotide may comprise at least one modified base moiety which is selected from the group including but not limited to 5-fluorouracil, 5-bromouracil,

- 30 5-chlorouracil, 5-iodouracil, hypoxanthine, xantine,
  4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil,
  5-carboxymethylaminomethyl-2-thiouridine,
  5-carboxymethylaminomethyluracil, dihydrouracil, beta-Dgalactosylqueosine, inosine, N6-isopentenyladenine,
- 35 1-methylguanine, 1-methylinosine, 2,2-dimethylguanine, 2-methyladenin, 2-methylguanine, 3-methylcytosin, 5-methylcytosine, N6-adenine, 7-methylguanine,

5-methylaminomethyluracil, 5-methoxyaminomethyl-2-thiouracil, beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil, 5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid (v), wybutoxosine, pseudouracil, queosine, 2-thiocytosin, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, uracil-5-oxyacetic acid methylester, uracil-5-oxyacetic acid (v), 5-methyl-2-thiouracil, 3-(3-amino-3-N-2-carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine.

The antisense oligonucleotide may also comprise at least one modified sugar moiety selected from the group including but not limited to arabinose, 2-fluoroarabinose, xylulose, and hexose.

In yet another embodiment, the antisense oligonucleotide

15 comprises at least one modified phosphate backbone selected
from the group consisting of a phosphorothioate, a
phosphorodithioate, a phosphoramidothioate, a
phosphoramidate, a phosphordiamidate, a methylphosphonate, an
alkyl phosphotriester, and a formacetal or analog thereof.

In yet another embodiment, the antisense oligonucleotide is an α-anomeric oligonucleotide. An α-anomeric oligonucleotide forms specific double-stranded hybrids with complementary RNA in which, contrary to the usual β-units, the strands run parallel to each other (Gautier et al., 1987, Nucl. Acids Res. 15:6625-6641). The oligonucleotide is a 2'-0-methylribonucleotide (Inoue et al., 1987, Nucl. Acids Res. 15:6131-6148), or a chimeric RNA-DNA analogue (Inoue et al., 1987, FEBS Lett. 215:327-330).

Oligonucleotides of the invention may be synthesized by standard methods known in the art, e.g. by use of an automated DNA synthesizer (such as are commercially available from Biosearch, Applied Biosystems, etc.). As examples, phosphorothicate oligonucleotides may be synthesized by the method of Stein et al. (1988, Nucl. Acids Res. 16:3209), standard pore glass polymer supports (Sarin et al., 1988, Proc. Natl. Acad. Sci. U.S.A. 85:7448-7451), etc.

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The antisens molecules should be delivered to cells which express the MC4-R in vivo, e.g., neural tissue. A number of methods have been developed for delivering antisense DNA or RNA to cells; e.g., antisense molecules can be injected directly into the tissue site, or modified antisense molecules, designed to target the desired cells (e.g., antisense linked to peptides or antibodies that specifically bind receptors or antigens expressed on the target cell surface) can be administered systemically.

- However, it is often difficult to achieve intracellular concentrations of the antisense sufficient to suppress translation of endogenous mRNAs. Therefore a preferred approach utilizes a recombinant DNA construct in which the antisense oligonucleotide is placed under the control of a strong pol III or pol II promoter. The use of such a
  - construct to transfect target cells in the patient will result in the transcription of sufficient amounts of single stranded RNAs that will form complementary base pairs with the endogenous MC4-R transcripts and thereby prevent
- 20 translation of the MC4-R mRNA. For example, a vector can be introduced in vivo such that it is taken up by a cell and directs the transcription of an antisense RNA. Such a vector can remain episomal or become chromosomally integrated, as long as it can be transcribed to produce the desired
- 25 antisense RNA. Such vectors can be constructed by recombinant DNA technology methods standard in the art. Vectors can be plasmid, viral, or others known in the art, used for replication and expression in mammalian cells. Expression of the sequence encoding the antisense RNA can be
- 30 by any promoter known in the art to act in mammalian, preferably human cells. Such promoters can be inducible or constitutive. Such promoters include but are not limited to: the SV40 early promoter region (Bernoist and Chambon, 1981, Nature 290:304-310), the promoter contained in the 3' long
- 35 terminal repeat of Rous sarcoma virus (Yamamoto et al., 1980, C ll 22:787-797), the herpes thymidine kinase promoter (Wagner et al., 1981, Proc. Natl. Acad. Sci. U.S.A. 78:1441-

1445), the regulatory sequences of the metallothionein gene (Brinster et al., 1982, Nature 296:39-42), etc. Any type of plasmid, cosmid, YAC or viral vector can be used to prepare the recombinant DNA construct which can be introduced

5 directly into the tissue site; e.g., the choroid plexus or hypothalamus. Alternatively, viral vectors can be used which selectively infect the desired tissue; (e.g., for brain, herpesvirus vectors may be used), in which case administration may be accomplished by another route (e.g., 10 systemically).

Ribozyme molecules designed to catalytically cleave MC4-R mRNA transcripts can also be used to prevent translation of MC4-R mRNA and expression of MC4-R. (See, e.g., PCT International Publication WO90/11364, published October 4, 15 1990; Sarver et al., 1990, Science 247:1222-1225). While ribozymes that cleave mRNA at site specific recognition sequences can be used to destroy MC4-R mRNAs, the use of hammerhead ribozymes is preferred. Hammerhead ribozymes cleave mRNAs at locations dictated by flanking regions that 20 form complementary base pairs with the target mRNA. requirement is that the target mRNA have the following sequence of two bases: 5'-UG-3'. The construction and production of hammerhead ribozymes is well known in the art and is described more fully in Haseloff and Gerlach, 1988, 25 Nature, 334:585-591. There are hundreds of potential hammerhead ribozyme cleavage sites within the nucleotide sequence of human MC4-R cDNA (see FIG. 5). Preferably the ribozyme is engineered so that the cleavage recognition site is located near the 5' end of the MC4-R mRNA; i.e., to 30 increase efficiency and minimize the intracellular accumulation of non-functional mRNA transcripts.

The ribozymes of the present invention also include RNA endoribonucleases (hereinafter "Cech-type ribozymes") such as the one which occurs naturally in Tetrahymena Thermophila

35 (known as the IVS, or L-19 IVS RNA) and which has been extensively described by Thomas Cech and collaborators (Zaug, et al., 1984, Science, 224:574-578; Zaug and Cech, 1986,

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Science, 231:470-475; Zaug, et al., 1986, Nature, 324:429-433; published International patent application No. WO 88/04300 by University Patents Inc.; Been and Cech, 1986, Cell, 47:207-216). The Cech-type ribozymes have an eight base pair active site which hybridizes to a target RNA sequence whereafter cleavage of the target RNA takes place. The invention encompasses those Cech-type ribozymes which target eight base-pair active site sequences that are present in MC4-R.

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As in the antisense approach, the ribozymes can be composed of modified oligonucleotides (e.g. for improved stability, targeting, etc.) and should be delivered to cells which express the MC4-R in vivo, e.g., hypothalamus. A preferred method of delivery involves using a DNA construct "encoding" the ribozyme under the control of a strong constitutive pol III or pol II promoter, so that transfected cells will produce sufficient quantities of the ribozyme to destroy endogenous MC4-R messages and inhibit translation. Because ribozymes unlike antisense molecules, are catalytic, a lower intracellular concentration is required for efficiency.

Endogenous MC4-r gene expression can also be reduced by inactivating or "knocking out" the MC4-r gene or its promoter using targeted homologous recombination (e.g., see Smithies 25 et al., 1985, Nature 317:230-234; Thomas & Capecchi, 1987, Cell 51:503-512; Thompson et al., 1989 Cell 5:313-321; each of which is incorporated by reference herein in its entirety). For example, a mutant, non-functional MC4-R (or a completely unrelated DNA sequence) flanked by DNA homologous 30 to the endogenous MC4-r gene can be used, with or without a selectable marker and/or a negative selectable marker, to transfect cells that express MC4-R in vivo. Insertion of the DNA construct, via targeted homologous recombination, results in inactivation of the MC4-r gene. Such approaches are 35 particularly suited in the agricultural field where modifications to ES (embryonic stem) cells can be used to generate animal offspring with an inactive MC4-R (e.g., see

Thomas & Capecchi 1987 and Thompson 1989, <u>supra</u>). However this approach can be adapted for use in humans provided the recombinant DNA constructs are directly administered or targeted to the required site <u>in vivo</u> using appropriate viral vectors, <u>e.g.</u>, herpes virus vectors for delivery to brain tissue; <u>e.g.</u>, the hypothalamus and/or choroid plexus.

Alternatively, endogenous MC4-R gene expression can be reduced by targeting deoxyribonucleotide sequences complementary to the regulatory region of the MC4-R gene

10 (i.e., the MC4-R promoter and/or enhancers) to form triple helical structures that prevent transcription of the MC4-R gene in target cells in the body. (See generally, Helene, C. 1991, Anticancer Drug Des., 6(6):569-84; Helene, C., et al., 1992, Ann, N.Y. Acad. Sci., 660:27-36; and Maher, L.J., 1992, 15 Bioassays 14(12):807-15).

## 5.4.3. DELIVERY OF SOLUBLE MC4-R POLYPEPTIDES

Genetically engineered cells that express soluble MC4-R ECDs or fusion proteins <u>e.g.</u> fusion Ig molecules can be

20 administered <u>in vivo</u> where they may function as "bioreactors" that deliver a supply of the soluble molecules. Such soluble MC4-R polypeptides and fusion proteins, when expressed at appropriate concentrations, should neutralize or "mop up" the native ligand for MC4-R, and thus act as inhibitors of MC4-R activity and induce weight gain.

# 5.5. PHARMACEUTICAL FORMULATIONS AND METHODS OF TREATING BODY WEIGHT DISORDERS

The invention encompasses methods and compositions for modifying body weight and treating body weight disorders, including but not limited to obesity, cachexia and anorexia. Because a loss of normal MC4-R gene product function results in the development of an obese phenotype, an increase in MC4-R gene product activity, or activation of the MC4-R pathway (e.g., downstream activation) would facilitate progress towards a normal body weight state in obese

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individuals exhibiting a deficient level of MC4-R gene expression and/or MC4-R activity.

Alternatively, symptoms of certain body weight disorders such as, for example, cachexia, which involve a lower than normal body weight phenotype, may be am liorated by decreasing the level of MC4-R gene expression, and/or MC4-R gene activity, and/or downregulating activity of the MC4-R pathway (e.g., by targeting downstream signalling events). Different approaches are discussed below.

Agonists of MC4-R can be used to induce weight loss for 10 treating obesity. Antagonists of MC4-R activity can be used to induce weight gain for treating conditions such as anorexia or cachexia. It is not necessary that the compound demonstrate absolute specificity for the MC4-R. For example, 15 compounds which agonize both MC4-R and MC1-R could be used; such compounds could be administered so that delivery to the brain is optimized to achieve weight reduction, and side effects, such as peripheral melanin production resulting in a "tan" may well be tolerated. Compounds which do not 20 demonstrate a specificity for MC4-R can be administered in conjunction with another therapy or drug to control the sideeffects that may result from modulating another melanocortin receptor; however, compounds which demonstrate a preference or selectivity for MC4-R over MC3-R are preferred since both 25 receptors are expressed in the brain where localized delivery cannot be used to compensate for lack of receptor specificity.

## 5.5.1. DOSE DETERMINATIONS

Toxicity and therapeutic efficacy of such compounds can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, e.g., for determining the LD<sub>50</sub> (the dose lethal to 50% of the population) and the ED<sub>50</sub> (the dose therapeutically effective in 50% of the population). The dose ratio between toxic and therapeutic effects is the therapeutic index and it can be expressed as

the ratio LD<sub>50</sub>/ED<sub>50</sub>. Compounds which exhibit large therapeutic

indices are preferred. While compounds that exhibit toxic side effects may be used, care should be taken to design a delivery system that targets such compounds to the site of affected tissue in order to minimize potential damage to 5 uninfected cells and, thereby, reduce side effects.

The data obtained from the cell culture assays and animal studies can be used in formulating a range of dosage for use in humans. The dosage of such compounds lies preferably within a range of circulating concentrations that 10 include the ED<sub>50</sub> with little or no toxicity. The dosage may vary within this range depending upon the dosage form employed and the route of administration utilized. For any compound used in the method of the invention, the therapeutically effective dose can be estimated initially 15 from cell culture assays. A dose may be formulated in animal models to achieve a circulating plasma concentration range that includes the  $IC_{50}$  (i.e., the concentration of the test compound which achieves a half-maximal inhibition of symptoms) as determined in cell culture. Such information 20 can be used to more accurately determine useful doses in humans. Levels in plasma may be measured, for example, by high performance liquid chromatography.

## 5.5.2. FORMULATIONS AND USE

Pharmaceutical compositions for use in accordance with the present invention may be formulated in conventional manner using one or more physiologically acceptable carriers or excipients.

Thus, the compounds and their physiologically acceptable 30 salts and solvates may be formulated for administration by inhalation or insufflation (either through the mouth or the nose) or oral, buccal, parenteral or rectal administration.

For oral administration, the pharmaceutical compositions may take the form of, for example, tablets or capsules

35 prepared by conventional means with pharmaceutically acceptable excipients such as binding agents (e.g., pregelatinised maize starch, polyvinylpyrrolidone or

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hydroxypropyl methylcellulose); fillers (e.g., lactose, microcrystalline cellulose or calcium hydrogen phosphate); lubricants (e.g., magnesium stearate, talc or silica); disintegrants (e.g., potato starch or sodium starch 5 glycolate); or wetting agents (e.g., sodium lauryl sulphate). The tablets may be coated by methods well known in the art. Liquid preparations for oral administration may take the form of, for example, solutions, syrups or suspensions, or they may be presented as a dry product for constitution with water 10 or other suitable vehicle before use. Such liquid preparations may be prepared by conventional means with pharmaceutically acceptable additives such as suspending agents (e.g., sorbitol syrup, cellulose derivatives or hydrogenated edible fats); emulsifying agents (e.g., lecithin 15 or acacia); non-aqueous vehicles (e.g., almond oil, oily esters, ethyl alcohol or fractionated vegetable oils); and preservatives (e.g., methyl or propyl-p-hydroxybenzoates or sorbic acid). The preparations may also contain buffer salts, flavoring, coloring and sweetening agents as 20 appropriate.

Preparations for oral administration may be suitably formulated to give controlled release of the active compound.

For buccal administration the compositions may take the form of tablets or lozenges formulated in conventional 25 manner.

For administration by inhalation, the compounds for use according to the present invention are conveniently delivered in the form of an aerosol spray presentation from pressurized packs or a nebulizer, with the use of a suitable propellant, 30 e.g., dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurized aerosol the dosage unit may be determined by providing a valve to deliver a metered amount. Capsules and cartridges of e.g. gelatin for use in 35 an inhaler or insufflator may be formulated containing a powder mix of the compound and a suitable powder bas such as lactose or starch.

The compounds may be formulated for parenteral administration by injection, e.g., by bolus injection or continuous infusion. Formulations for injection may be presented in unit dosage form, e.g., in ampoules or in multi-5 dose containers, with an added preservative. The compositions may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents. Alternatively, the active ingredient may 10 be in powder form for constitution with a suitable vehicle, e.g., sterile pyrogen-free water, before use.

The compounds may also be formulated in rectal compositions such as suppositories or retention enemas, <u>e.g.</u>, containing conventional suppository bases such as cocoa butter or other glycerides.

In addition to the formulations described previously, the compounds may also be formulated as a depot preparation. Such long acting formulations may be administered by implantation (for example subcutaneously or intramuscularly) or by intramuscular injection. Thus, for example, the compounds may be formulated with suitable polymeric or hydrophobic materials (for example as an emulsion in an acceptable oil) or ion exchange resins, or as sparingly soluble derivatives, for example, as a sparingly soluble 25 salt.

The compositions may, if desired, be presented in a pack or dispenser device which may contain one or more unit dosage forms containing the active ingredient. The pack may for example comprise metal or plastic foil, such as a blister pack. The pack or dispenser device may be accompanied by instructions for administration.

# 5.6. <u>DIAGNOSIS OF BODY WEIGHT DISORDER ABNORMALITIES</u> Mutations at a number of different genetic loci have

35 been identified which lead to phenotypes related to body weight disorders. Ideally, the treatment of patients suffering from such body weight disorders will be designed to

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target the particular genetic loci containing the mutation.

Therefore, diagnostic methods that identify mutations in specific genes related to body weight disorders, such as the MC4-r gene, will permit the treatment of body weight disorders through targeting of the mutated gene.

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A variety of methods can be employed for the diagnostic and prognostic evaluation of body weight disorders, including obesity, cachexia and anorexia, and for the identification of subjects having a predisposition to such disorders.

- Such methods may, for example, utilize reagents such as the MC4-r gene nucleotide sequences and antibodies directed against MC4-r gene products, including peptide fragments thereof. Specifically, such reagents may be used, for example, for: (1) the detection of the presence of MC4-r gene mutations, or the detection of either over- or under-expression of MC4-r gene mRNA relative to the non-body weight disorder state; and (2) the detection of either an over- or an under-abundance of MC4-r gene product relative to the non-body weight disorder state.
- The methods described herein may be performed, for example, by utilizing pre-packaged diagnostic kits comprising at least one specific MC4-r gene nucleic acid or anti-MC4-r antibody reagent, which may be conveniently used, e.g., in clinical settings, to diagnose patients exhibiting body weight disorder abnormalities.

For the detection of MC4-r mutations, any nucleated cell can be used as a starting source for genomic nucleic acid. For the detection of MC4-r gene expression or MC4-r gene products, any cell type or tissue in which the MC4-r gene is expressed, such as, for example, brain cells, may be utilized.

Nucleic acid-based detection techniques are described, below, in Section 5.6.1. Peptide detection techniques are described, below, in Section 5.6.2.

5.6.1. <u>DETECTION OF MC4-r GENE NUCLEIC ACID MOLECULES</u>
Predisposition to body weight disorders can be
ascertained by testing any tissue for mutations of the MC4-r
gene. For example, a person who has inherited a germline
5 MC4-r mutation would be more likely to develop an obese
phenotype. In addition, prenatal diagnosis can be carried
out by testing fetal cells, placental cells or amniotic fluid
for mutations of the MC4-r gene. Alterations in the MC4-r
allele can be detected using any of the methods discussed
10 herein. Mutations within the MC4-r gene can be detected by

Nucleic acid from any nucleated cell can be used as the starting point for such assay techniques, and may be isolated according to standard nucleic acid preparation procedures

15 which are well known to those of skill in the art.

utilizing a number of techniques.

DNA may be used in hybridization or amplification assays of biological samples to detect abnormalities involving MC4-r gene structure, including point mutations, insertions, deletions and chromosomal rearrangements. Such assays may 20 include, but are not limited to, Southern analyses, single stranded conformational polymorphism analyses (SSCP), and PCR analyses.

Techniques well-known in the art may be used to scan regions of a selected genome or portion thereof, for any

25 variant or mutation, be it known or unknown. Methods that may be used to detect such variants or mutations in the MC4-r gene include direct sequencing (Maxam & Gilbert, 1980, Methods in Enzymology 65:499-560; Sanger et al., 1977, Proc. Natl. Acad. Sci. 74:5463); resequencing by oligonucleotide

30 arrays (Fodor et al., 1993, Nature 364:555-556); Southern blot or pulsed-field gel (PFGE) analysis (Schwartz et al., 1984, Cell 37:67); single stranded conformation analysis ("SSCA") (Orita et al., 1989, Proc. Natl. Acad. Sci. USA 86:2776-2770); heteroduplex analysis (HA) (Keen et al., 1991, Trends Genet. 7:5); denaturing gradient gel electrophoresis ("DGGE") (Wartell et al., 1990, Nucl. Acids Res. 18:2699-2705; Myers et al., 1985, Nucleic Acid Research 13:3131-

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3145); denaturing HPLC (Underhill, P.A., 1997, Proc. Natl. Acad. Sci USA 93:196-200); RNase protection assays (Finkelstein et al., 1990, Genotics 7:167-172; Kinsler et al., 1991, Science 251:1366-1370; Myers, R.M. et al., 1985, 5 Science 230:1242-1246); allele specific oligonucleotide ("ASO") hybridization (Conner et al., 1983, Proc. Natl. Acad. Sci. 80:278-282; Wallace, R.B., et al., 1979, Nucl. Acids Res. 6:3543-3557; Saiki, R.K. et al., 1989, Proc. Natl. Acad. Sci. USA 86:6230-6234)); oligonucleotide ligation assay 10 ("OLA") (Landegren, U., et al., 1988, Science 241:1077-1080; Tobe, V.O., et al., 1996, Nucl. Acids Res. 24:3728-3732); sequence-specific amplification (Newton, C.R. et al., 1989, Nucl. Acids Res. 17:2503-2516); chemical mismatch cleavage (CMC) (Cotton, R.G.H. et al., 1988, Nucl. Acids Res. 17:4223-15 4233); enzymatic mismatch cleavage (EMC) (Babon J.J., R. et al., 1995, Nucl. Acids Res. 23:5082-5084; Marshall, R.D. et al., 1995, Nat. Genet. 9:177-183); and the use of proteins which recognize nucleotide mismatches, such as the E. coli mutS protein (Modrich, 1991, Ann. Rev. Genet. 25:229-253). Diagnostic methods that may be used to detect mutations 20 at the MC4-r locus can classified into two general categories of use. The first category of diagnostic methods includes

at the MC4-r locus can classified into two general categories of use. The first category of diagnostic methods includes those diagnostic methods designed to scan a region for an unknown variant. These diagnostic methods may also be applied to the detection of known variants. The second category of diagnostic methods includes those designed specifically to type previously-identified variants, but are not usually applied to the detection of unknown variants.

Diagnostic methods that may be used to type previously30 identified variants may include, but are not limited to,
allele-specific oligonucleotide (ASO) hybridization, and
oligonucleotide ligation assay (OLA), and sequence-specific
amplification.

In one embodiment, allele-specific oligonucleotide 35 hybridization is used to detect a previously-identified variant(s) or allele(s) of MC4-R. Allele-specific oligonucleotide hybridization comprises the separate hybridization of a pair of oligonucleotides, specific to the previously-identified allele, to PCR-amplified genomic DNA or RNA, under conditions that discriminate between complete matches and single-base mismatches. The pair of oligonucleotides must encompass the variant base. Under the

5 oligonucleotides must encompass the variant base. Under the appropriate reaction conditions, the target DNA is not amplified if there is a base mismatch (e.g., a nucleotide substitution caused by a mutation) or a small deletion or insertion, at the 3' end of the primer (Okayama et al, 1989,

10 J. Lab. Clin. Med. 114:105-113; Sommer et al., 1992, BioTechniques 12:82-87).

In a specific embodiment of the invention the following oligonucleotides, or their complement, may be used to identify MC4-R varients. To identify the Ile137Thr mutation the following oligonucleotides may be used:

- 5'-CACTT-3'
- 5'-ATCCACTTGC-3'
- 5'-TGCATCCACTTGCAG-3'
- 5'-GCTTGCATCCACTTGCAGCC-3'
- 20 5'-CTTGCTTGCATCCACTTGCAGCCTG-3'
  - 5'-CTCCTTGCTTGCATCCACTTGCAGCCTGCT-3'.

To identify the Vallo2Ile mutation the following oligonucleotides may be used:

- 5'-CCATT-3'
- 25 5'-AAACCATTAT-3'
  - 5'-CAGAAACCATTATCA-3'
  - 5'-GATCAGAAACCATTATCATC-3'
  - 5'-ATGGATCAGAAACCATTATCATCAC-3'
  - 5'-CAAATGGATCAGAAACCATTATCATCACCC-3'.
- 30 To identify the Thr112Met mutation the following oligonucleotides may be used;
  - 5'-TATGG-3'
  - 5'-AGATATGGAT-3'
  - 5'-TACAGATATGGATGC-3'
- 35 5'-CAGTACAGATATGGATGCAC-3'
  - 5'-GTACAGTACAGATATGGATGCACAG-3'
  - 5'-ACAGTACAGTACAGATATGGATGCACAGAG-3'.

In another embodiment, an oligonucleotide ligation assay (OLA) is used to discriminate a single base mismatch, through the use of a pair of oligonucleotides that are complementary to two preselected alleles. How ver, in an oligonucleotide

- 5 ligation assay, the two oligonucleotides of the pair are ligated in a template-dependent fashion at the site of the variant base. The template or target sequence is typically a region of the gene encompassing the variant base, and is preferably PCR-amplified from genomic DNA or RNA. Similarly,
- 10 as described above, sequence-specific amplification relies on PCR primers designed such that the variant base is located at the 3' end of one of the PCR primers. Thus PCR amplification is dependent on complete annealing of this primer and will not take place if there exists a single-base mismatch.
- In specific embodiments, the direct sequencing of PCR products that are amplified from genomic DNA or RNA may be accomplished using either fluorescent or radioactive methods well known to those skilled in the art.

In another embodiment, resequencing of these PCR

20 products is accomplished more rapidly by hybridization, to high-density arrays, of oligonucleotides representing the wild-type sequence of the MC4-R gene (Hattori, M., 1993, Genomics 15:415-417).

In another embodiment, Southern blot pulsed-field gel 25 (PFGE) analysis can be used to identify those sequence variants that abolish or create novel restriction sites.

In another embodiment, variants are identified by single-stranded conformational analysis (SSCA) by virtue of their effect on the conformation of the DNA molecule.

- 30 Regions containing known variants or regions being searched for variants are amplified by PCR. These products are denatured and electrophoresed through polyacrylamide gels under non-denaturing conditions that allow the single-stranded molecule to retain their intrastrand interactions.
- 35 Single-base changes affect these interactions and consequently alter the rate of migration of the molecule

through the gel. Thus, variants are identified as fragments of altered mobility.

In another embodiment, a SSCA related method of heteroduplex analysis detects single base-pair mismatches in 5 the sequences of re-annealed double-stranded molecules by detecting the altered conformations of such molecules. Such conformational changes will affect the migration of the reannealed double-stranded molecules through polyacrylamide gels under non-denaturing conditions.

electrophoresis (DGGE) may be used to identify variants based on the difference in melting temperature between two DNA fragments with a single base-pair difference. Double-stranded molecules are electrophoresed through a polyacrylamide gel containing a gradient of increasing denaturant, e.g. example of denaturant, such that at a defined point, a molecule will begin to denature and migrate more slowly. As with SSCA and HA, variants are identified by their altered mobility in the gel.

In another embodiment, denaturing HPLC may be used to reveal variants that alter the conformation of the DNA fragment and thus affect migration rate through chromatographic columns. Some methods of denaturing HPLC involve denaturing and annealing a test DNA or RNA to a control to allow heteroduplexes to form, as in HA analysis. Subsequently chemicals or enzymes are used to cleave one or two strands at or nearby the mismatch. In one embodiment, chemical mismatch cleavage (CMC), which employs osmium tetroxide, hydroxylamine and piperidine, is used to cleave one or both strands.

In another embodiment, enzymatic mismatch cleavage (EMC), using mismatch-cleavage enzymes such as T4 endonuclease VII, is used to cleave one or both strands.

In another embodiment, RNase protection assays are used 35 to detect an RNA:RNA mismatch. RNase protection assays exploit the ability of RNase A to cleave RNA:RNA mismatches. Amplified DNA from genomic DNA or RNA is reverse transcribed

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into cRNA. The test sample cRNA is annealed to a control wild-type cRNA. Sequence variants may then be detected as mismatches between the two cRNA molecules that are cleaved by RNase. Cleavage is revealed by electrophoretic sizing of the 5 products.

Among the MC4-r nucleic acid sequences which are preferred for such hybridization and/or PCR analyses are those which will detect the presence of the MC4-r gene mutations described, below, in Section 8.2.

The level of MC4-r gene expression can also be assayed. For example, RNA from a cell type or tissue known, or suspected, to express the MC4-r gene, such as brain, may be isolated and tested utilizing hybridization or PCR techniques such as are described, above. The isolated cells can be derived from cell culture or from a patient. The analysis of cells taken from culture may be a necessary step in the assessment of cells to be used as part of a cell-based gene therapy technique or, alternatively, to test the effect of compounds on the expression of the MC4-r gene. Such analyses may reveal both quantitative and qualitative aspects of the expression pattern of the MC4-r gene, including activation or inactivation of MC4-r gene expression.

In one embodiment of such a detection scheme, a cDNA molecule is synthesized from an RNA molecule of interest

25 (e.g., by reverse transcription of the RNA molecule into cDNA). A sequence within the cDNA is then used as the template for a nucleic acid amplification reaction, such as a PCR amplification reaction, or the like. The nucleic acid reagents used as synthesis initiation reagents (e.g.,

30 primers) in the reverse transcription and nucleic acid amplification steps of this method are chosen from among the MC4-r gene nucleic acid reagents described in Section 5.1. The preferred lengths of such nucleic acid reagents are at least 9-30 nucleotides. For detection of the amplified

35 product, the nucleic acid amplification may be performed using radioactively or non-radioactively labeled nucleotides. Alternatively, enough amplified product may be made such that

the product may be visualized by standard ethidium bromide staining or by utilizing any other suitable nucleic acid staining method.

Additionally, it is possible to perform such MC4-r gene
5 expression assays "in situ", <u>i.e.</u>, directly upon tissue
sections (fixed and/or frozen) of patient tissue obtained
from biopsies or resections, such that no nucleic acid
purification is necessary. Nucleic acid reagents such as
those described in Section 5.1 may be used as probes and/or
10 primers for such in situ procedures (see, for example, Nuovo,
G.J., 1992, "PCR In Situ Hybridization: Protocols And
Applications", Raven Press, NY).

Alternatively, if a sufficient quantity of the appropriate cells can be obtained, standard Northern analysis 15 can be performed to determine the level of mRNA expression of the MC4-r gene.

## 5.6.2. <u>DETECTION OF MC4-r GENE PRODUCTS</u>

Antibodies directed against wild type or mutant MC4-r

20 gene products or conserved variants or peptide fragments
thereof, may also be used as body weight disorder diagnostics
and prognostics, as described herein. Such diagnostic
methods, may be used to detect abnormalities in the level of
MC4-r gene expression, or abnormalities in the structure

25 and/or temporal, tissue, cellular, or subcellular location of
MC4-r gene product. Antibodies, or fragments of antibodies,
such as those described below, may be used to screen
potentially therapeutic compounds in vitro to determine their
effects on MC4-r gene expression and MC4-r peptide

30 production. The compounds which have beneficial effects on body weight disorders, such as obesity, cachexia and anorexia, can be identified, and a therapeutically effective dose determined.

In vitro immunoassays may also be used, for example, to 35 assess the efficacy of cell-based gene therapy for body weight disorders, including obesity, cachexia and anorexia. Antibodies directed against MC4-r peptides may be used in

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vitro to determine the lev l of MC4-r gene expression achieved in cells genetically engineered to produce MC4-r peptides. Such analysis will allow for a determination of the number of transformed cells necessary to achieve 5 therapeutic efficacy in vivo, as well as optimization of the gene replacement protocol.

The tissue or cell type to be analyzed will generally include those which are known, or suspected, to express the MC4-r gene, such as, for example, brain cells. The protein isolation methods employed herein may, for example, be such as those described in Harlow and Lane (Harlow, E. and Lane, D., 1988, "Antibodies: A Laboratory Manual", Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York), which is incorporated herein by reference in its entirety. The isolated cells can be derived from cell culture or from a patient. The analysis of cell taken from culture may be a necessary step in the assessment of cells to be used as part of a cell-based gene therapy technique or, alternatively, to test the effect of compounds on the expression of the MC4-r 20 gene.

Preferred diagnostic methods for the detection of wildtype or mutant MC4-r gene products or conserved variants or
peptide fragments thereof, may involve, for example,
immunoassays wherein the MC4-r gene products or conserved
25 variants or peptide fragments are detected by their
interaction with an anti-MC4-r gene product-specific
antibody.

For example, antibodies, or fragments of antibodies, such as those described, above, in Section 5.3, useful in the 30 present invention may be used to quantitatively or qualitatively detect the presence of wild type or mutant MC4-r gene products or conserved variants or peptide fragments thereof. This can be accomplished, for example, by immunofluorescence techniques employing a fluorescently 35 labeled antibody (see below, this Section) coupled with light microscopic, flow cytometric, or fluorimetric detection.

Such techniques are especially preferred if such MC4-r gene products are expressed on the cell surface.

The antibodies (or fragments thereof) useful in the present invention may, additionally, be employed 5 histologically, as in immunofluorescence or immunoelectron microscopy, for in situ detection of wild type or mutant MC4r gene products or conserved variants or peptide fragments thereof. In situ detection may be accomplished by removing a histological specimen from a patient, and applying thereto a 10 labeled antibody of the present invention. The antibody (or fragment) is preferably applied by overlaying the labeled antibody (or fragment) onto a biological sample. Through the use of such a procedure, it is possible to determine not only the presence of the MC4-r gene product, or conserved variants 15 or peptide fragments, but also its distribution in the examined tissue. Using the present invention, those of ordinary skill will readily perceive that any of a wide variety of histological methods (such as staining procedures) can be modified in order to achieve such in situ detection.

Immunoassays for wild type or mutant MC4-r gene products or conserved variants or peptide fragments thereof will typically comprise incubating a sample, such as a biological fluid, a tissue extract, freshly harvested cells, or lysates of cells which have been incubated in cell culture, in the presence of a detectably labeled antibody capable of identifying wild type or mutant MC4-r gene products or conserved variants or peptide fragments thereof, and detecting the bound antibody by any of a number of techniques well-known in the art.

The biological sample may be brought in contact with and immobilized onto a solid phase support or carrier such as nitrocellulose, or other solid support which is capable of immobilizing cells, cell particles or soluble proteins. The support may then be washed with suitable buffers followed by treatment with the detectably labeled MC4-r gene specific antibody. The solid phase support may then be washed with the buffer a second time to remove unbound antibody. The

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amount of bound label on solid support may then be detected by conventional means.

By "solid phase support or carrier" is intended any support capable of binding an antigen or an antibody. Well-5 known supports or carriers include glass, polystyrene, polypropylene, polyethylene, dextran, nylon, amylases, natural and modified celluloses, polyacrylamides, gabbros, and magnetite. The nature of the carrier can be either soluble to some extent or insoluble for the purposes of the 10 present invention. The support material may have virtually any possible structural configuration so long as the coupled molecule is capable of binding to an antigen or antibody. Thus, the support configuration may be spherical, as in a bead, or cylindrical, as in the inside surface of a test 15 tube, or the external surface of a rod. Alternatively, the surface may be flat such as a sheet, test strip, etc. Preferred supports include polystyrene beads. Those skilled in the art will know many other suitable carriers for binding antibody or antigen, or will be able to ascertain the same by 20 use of routine experimentation.

The binding activity of a given lot of anti-MC4-r gene product antibody may be determined according to well known methods. Those skilled in the art will be able to determine operative and optimal assay conditions for each determination by employing routine experimentation.

One of the ways in which the MC4-r gene peptide-specific antibody can be detectably labeled is by linking the same to an enzyme and use in an enzyme immunoassay (EIA) (Voller, A., "The Enzyme Linked Immunosorbent Assay (ELISA)", 1978,

- Ouarterly Publication, Walkersville, MD); Voller, A. et al., 1978, J. Clin. Pathol. 31:507-520; Butler, J.E., 1981, Meth. Enzymol. 73:482-523; Maggio, E. (ed.), 1980, Enzyme Immunoassay, CRC Press, Boca Raton, FL,; Ishikawa, E. et al.,
- 35 (eds.), 1981, Enzyme Immunoassay, Kgaku Shoin, Tokyo). The enzyme which is bound to the antibody will react with an appropriate substrate, preferably a chromogenic substrate, in

such a manner as to produce a chemical moi ty which can be detected, for example, by spectrophotometric, fluorimetric or Enzymes which can be used to detectably by visual means. label the antibody include, but are not limited to, malate 5 dehydrogenase, staphylococcal nuclease, d lta-5-steroid isomerase, yeast alcohol dehydrogenase, alphaglycerophosphate, dehydrogenase, triose phosphate isomerase, horseradish peroxidase, alkaline phosphatase, asparaginase, glucose oxidase, beta-galactosidase, ribonuclease, urease, 10 catalase, glucose-6-phosphate dehydrogenase, glucoamylase and acetylcholinesterase. The detection can be accomplished by colorimetric methods which employ a chromogenic substrate for the enzyme. Detection may also be accomplished by visual comparison of the extent of enzymatic reaction of a substrate 15 in comparison with similarly prepared standards.

Detection may also be accomplished using any of a variety of other immunoassays. For example, by radioactively labeling the antibodies or antibody fragments, it is possible to detect MC4-r gene peptides through the use of a 20 radioimmunoassay (RIA) (see, for example, Weintraub, B., Principles of Radioimmunoassays, Seventh Training Course on Radioligand Assay Techniques, The Endocrine Society, March, 1986, which is incorporated by reference herein). The radioactive isotope can be detected by such means as the use 25 of a gamma counter or a scintillation counter or by

It is also possible to label the antibody with a fluorescent compound. When the fluorescently labeled antibody is exposed to light of the proper wave length, its 30 presence can then be detected due to fluorescence. Among the most commonly used fluorescent labeling compounds are fluorescein isothiocyanate, rhodamine, phycoerythrin, phycocyanin, allophycocyanin, o-phthaldehyde and fluorescamine.

autoradiography.

35 The antibody can also be detectably labeled using fluorescence emitting metals such as <sup>152</sup>Eu, or others of the lanthanide series. These metals can be attached to the

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antibody using such metal chelating groups as diethylenetriaminepentacetic acid (DTPA) or ethylenediaminetetraacetic acid (EDTA).

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The antibody also can be detectably labeled by coupling 5 it to a chemiluminescent compound. The presence of the chemiluminescent-tagged antibody is then determined by detecting the presence of luminescence that arises during the course of a chemical reaction. Examples of particularly useful chemiluminescent labeling compounds are luminol, 10 isoluminol, theromatic acridinium ester, imidazole, acridinium salt and oxalate ester.

Likewise, a bioluminescent compound may be used to label the antibody of the present invention. Bioluminescence is a type of chemiluminescence found in biological systems in,

15 which a catalytic protein increases the efficiency of the chemiluminescent reaction. The presence of a bioluminescent protein is determined by detecting the presence of luminescence. Important bioluminescent compounds for purposes of labeling are luciferin, luciferase and aequorin.

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# 6. EXAMPLE: GENERATION OF AN MC4-R DEFICIENT MOUSE The following example describes the engineering and generation of "knock-out" mice in which the endogenous MC4-r is inactivated. The results show that the knock-out mice 25 gain weight, thus, demonstrating the role and function of the

MC4-R in body weight regulation.

## 6.1. MATERIALS AND METHODS

## 6.1.1. IDENTIFICATION OF THE MURINE MC4-R GENE

The murine melanocortin 4 receptor (MC4-r) gene was isolated from a mouse strain 129/Sv genomic phage library, obtained from Stratagene, using a human MC4-r probe. The human probe was generated by PCR amplification of MC4-r coding sequences from human genomic DNA using the following primers:

5'-ATA GTC GAC ATG GTG AAC TCC ACC CAC CGT-3'; and 5'-TAT AAG CTT TTA ATA TCT GCT AGA CAA GTC-3'.

Two positive phage clones containing the MC4-r gene were identified, and the MC4-r locus was subcloned from phage into pBluescript II as an ~5 Kb Hind III fragment, and an ~4.7 Kb Sac I fragment. These subclones were restriction mapped and partially sequenced to produce the map of the MC4-r locus shown in Fig. 2A. In order to inactivate MC4-r, a targeting construct was built which would delete the majority of MC4-r coding sequences following homologous recombination with the endogenous MC4-r locus.

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## 6.1.2. GENERATION OF THE TARGETING CONSTRUCT

The MC4-r targeting construct was constructed in the following manner. The 1.4 Kb Eco RI-Ava I fragment of pBR322 was replaced with the following synthetic oligonucleotides:

15 5'-AAT TAG CGG CCG CAG TAT GCA AAA AAA AGC CCG CTC ATT AGG CGG GCT-3'; and 5'-CCG AAG CCC GCC TAA TGA GCG GGC TTT TTT TTG CAT ACT GCG GCC GCT-3'.

The resulting plasmid, called pJN1, was digested with Not I and the following oligonucleotides were ligated into the Not 20 I site.

5'-GGC CGG CAT GCA TCA AGC TTA TCT CGA GAT CGT CGA CTA CCA
TGG TAC ATC GAT CAG GTA CCA TCC CGG GGC-3'; and 5'GGC CGC
CCC GGG ATG GTA CCT GAT CGA TGT ACC ATG GTA GTC GAC GAT CTC
GAG ATA AGC TTG ATG CAT GCC-3'.

The resulting plasmid was called pJN2.

The 1.2 Kb Sph I-Hind III fragment 3'of the MC4-r gene (see Fig. 2A) was subcloned into SphI-Hind III digested pJN2 to generate the plasmid MC4-r KO 3' (FIG. 2B). This fragment represents the 3' region of genomic homology in the targeting vector. A 3.4 Kb NcoI-Hind III fragment, including the first approximately 20 nucleotides of the MC4-r gene (see Fig. 2A), was excised as a NcoI-Asp718 fragment from the subclone MC4-r locus. The Asp718 site was derived from pBluescript II polylinker sequences immediately flanking the native Hind III site approximately 3.4 Kb 5' of the MC4-r gene (Fig. 2A). This fragment, which r presents the 5' region of genomic homology in the targeting construct, was ligated into NcoI-

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Asp 718 digested MC4-r KO 5' to produce MC4-r KO 5'3' (Fig. 2C).

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The PGK-neo expression cassette from the plasmid pKJ1 (Tybulewicz et al., Cell 65, 1153-1163, 1991), containing the 5 neo gene under the transcriptional control of the mouse phosphoglycerate kinase (PGK-1) promoter and the PGK-1 poly(A) addition site, was subcloned as an Eco RI-Hind III fragment into EcoRI-Hind III digested pGEM 7-Zf(+) to generate pGEM 7 (KJ1). The 1.7 Kb fragment containing the 10 PGK-neo expression cassette was excised by: 1) digestion of pGEM 7 (KJ1) with Xho I, which cuts in the polylinker 5' of the PGK promoter, and blunt end filling of the Xho I site with Klenow polymerase, and 2) digestion with Sca I which cuts within mouse genomic sequence 3' of the PGK 15 polyadenylation signal. This fragment was ligated into Xho I digested MC4-r KO 5'3' which had also been blunt-ended with Klenow polymerase, to generate the MC4-r targeting vector MC4-r KO 5'3' neo (Fig. 2D). A schematic map of the gene targeting strategy for inactivation of the MC4-r locus with 20 this vector is shown in Figures 3A-3D.

## 6.1.3. GENERATION OF TARGETED ES CELLS

The RF-8 ES cell line (obtained from the Gladstone Institute of Cardiovascular Disease, UCSF) was cultured on 25 SNL76/7 mitotically inactive feeder cells as described in McMahon and Bradley (1990, Cell 62: 1073-1085). For electroporation, cells were trypsinized and resuspended at a concentration of 1.1 x 10<sup>7</sup>/ml in PBS (Ca<sup>2+</sup> and MG<sup>2+</sup> free; Gibco). An 0.9 ml aliquot (1 x 10<sup>7</sup> cells) was mixed with 20 μg of MC4-r KO 5'3' DNA, which had been linearized by Not I digestion, and pulsed at 250V, 500 μF (Bio-Rad Gene Pulser), after which the cells were diluted in culture medium, plated at 1 x 10<sup>6</sup> per 100 mm plate containing feeder cells, and placed under selection twenty-four hours later in G418 sulfate (400 μg/ml powder, Gibco) for 6 days. 427 G418 r sistant clones were picked, dissociated with trypsin and divided into one well each of two 96-well plates. Upon

confluence, ES cells were frozen in one of the 96-well plates as described by Ramirez-Solis et al., (Methods in Enzymology, vol. 225, Wassarman, P.M., DePamphilis, M.L. (eds). Academic Press, p. 855-878, 1992) and expanded into a 24 well plate.
5 Upon confluence, DNA was prepared for Southern blot analysis.

Genomic DNA was prepared in situ from ES cells in 24 well plates by the procedure of Laird et al. (1991, Nucleic Acids Research 19: 4293). To screen for homologous recombination between the vector and the endogenous MC4-r 10 locus approximately 20 μg of genomic DNA was digested with Apa I, electrophoresed through a 1% agarose gel, transferred to Hybond N° membrane (Amersham), and hybridized with the <sup>32</sup>P radiolabeled Sac I-Sph I probe (see FIG. 3A).

## 6.1.4. GENERATION OF MC4-R DEFICIENT MICE

Clone 155 was injected into C57BL/6J blastocysts to generate chimeric mice as described in (Bradley, A. In Robertson, E.J. (ed) Teratocarcinomas and Embryonic Stem Cells. IRL Press, Oxford, England, p. 113-151, 1987). Male

- offspring (representing germline transmission of the ES genome) were screened for the presence of the targeted MC4-r gene by Southern blot hybridization of Apa I as well as Nco I digested tail DNA using the probe shown in FIG. 3A.
- 25 Offspring heterozygous for the mutation were identified by either the presence of a 7.6 Kb Apa I band in addition to the wild type 2.2 Kb band or the presence of a 1.9 Kb Nco I band in addition to the wild type bands of 2.6 Kb and 2.8 Kb (FIG. 3D).
- 30 Heterozygous mice were interbred and offspring generated by these matings were screened by Southern blot hybridization of Apa I as well as Nco I digested tail DNA. Mice homozygous for the deleted MC4-r gene were identified by the absence of the wild type 2.2 Kb Apa I band and 2.8 Kb Nco I band, and
- 35 presence of the targeted 7.6 Kb Apa I band and 1.9 Kb Nco I band. To verify deletion of the MC4-r gene, the Apa I digested and Nco I digested blots were stripped and re-probed

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with the human MC4-r coding sequence. No hybridizing bands were observed in the DNA from mice homozygous for the MC4-r mutation, verifying the absence of the MC4-r gene in these mice.

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#### 6.1.4 WEIGHT AND LENGTH MEASUREMENTS

Weight gain was regularly measured, beginning at 3-4 weeks of age, using a Sartorius model #14800 P balance.

Length was measured by manual immobilization and extension of the mouse to its full length, always by the same individual, and measurement of the nose to anus distance in centimeters.

#### 6.1.5. FOOD CONSUMPTION

Food intake was measured for two A<sup>y</sup>, two C57BL/6J, four
15 MC4-R homozygous mutants, and four wild type F2 controls,
each housed two to a cage. The mice were housed for at least
a week before any measurements were taken. Over a two week
period, a sufficient amount of food for the week was then
weighed and provided to the mice ad libitum. Each weekday
20 morning the remaining food was measured, for a total of eight
measurements. Cages were carefully monitored for spillage,
which was negligible. The A<sup>y</sup> and C57BL/6J mice were 9 weeks
of age at the time measurement of food consumption was
initiated; both the four MC4-R deficient mice and the F2
25 controls were each 15, 15, 17.5 and 20.5 weeks of age.

#### 6.1.6. SERUM ANALYSIS

For glucose, insulin and leptin measurements, blood was collected by retroorbital sinus puncture from animals

30 provided with food and water ad libitum. Mice were handled regularly (three times per week for several weeks) prior to bleeding to minimize stress, and cages were singly moved to a separation location at the time of bleeding. For measurement of glucose levels, 5 µl of serum was analyzed in a YSI

35 Model 27 glucose analyzer (Yellow Springs Instrument Company, Inc.) using a glucose oxidase assay. Results are expressed as mg/dl. The range of detection if 0-500 mg/dl, with a

coefficient of variation of <1%. Serum insulin concentration was measured in duplicate in a 10  $\mu$ l volume by a specific competitive protein binding assay using rat insulin as the standard. Results are expressed as ng/ml. The range of 5 detection is 0.1-25 ng/ml with a coefficient of variation of <10%. Leptin was measured in duplicate in 20  $\mu$ l of serum using a radioimmunoassay kit to mouse leptin with recombinant mouse leptin as the standard (Linco Research, Inc.).

For serum corticosterone measurements, mice were housed 10 at three animals per cage with food and water ad libitum. prevent stress-mediated elevation of corticosterone levels mice were handled 2-3 times/day for three days prior to drawing blood. Cages were brought one at a time into a separate room, mice were weighed and then held as if blood 15 were to be drawn. On the fourth day mice were handled similarly, and blood drawn between 8:00 and 9:00 A.M. within ' 30 sec of handling. Cages were not returned to the housing room until all the samples had been obtained. Blood was obtained by snipping the tail tip and collecting blood into a 20 Multivette S Gel tube (Sarstedt). Tubes were placed on ice for 20-40 minutes and centrifuged 3-4 minutes at 14,000 rpm to separate the serum. Two one  $\mu$ l aliquots of serum from each sample were then assayed for corticosterone levels using an ImmuChem Double Antibody Corticosterone 125I RIA kit (ICN 25 Biomedicals, Inc.)

#### 6.1.7. <u>HISTOLOGY</u>

For in situ hybridization analysis of POMC gene expression, wild type, heterozygous, and homozygous mutant 30 mice were maintained under a 12-h light, 12-h dark cycle at constant temperature. Food (Purina mouse chow) and water were provided ad libitum. Anesthetized (avertin) animals were sacrificed between 1500 and 1700 hrs. before lights out via cardiac puncture and perfusion with saline (20 mls) and 35 then 50 mls of ice-cold fixation buffer (4% paraformaldehyde in borate buffer, pH 9.5). Whole brains were rapidly removed and then post-fixed overnight in 10% sucrose/fixative buffer.

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Blocked hypothalamic sections wer frozen in powdered dry ice and then stored at -80°C prior to sectioning.

Antisense POMC probe was prepared by linearizing the plasmid mPOMCE3ribo (kindly provided by Dr. Malcolm Low),

5 containing exon 3 of the mouse POMCd gene, with Nco I. [35S] cRNA probes were prepared by transcribing 1 µg of each linearized DNA with T7 DNA polymerase for 1 hr at 37°C as described (Promega). Hypothalamic brain blocks were mounted on a frozen stage and serially sectioned into 4 series of

10 20-µM slices with a sliding microtome. Sections were prepared and hybridized for 20 h at 58°C with 35S-labeled probes (5 x 106 cpm/ml) in 65% formamide, 0.26 M NaCl, 1.3 x Denhardt's solution, 1.3 mM EDTA, 13% dextran sulfate, 13 mM Tris, pH 8. Sections were then digested with RNase

15 (20 µg/ml) for 30 min at 37°C, and then desalted in a series of washes from 4 x SSC/1 mM DTT to a final stringency of 0.1 x SSC/1 mM DTT at 65°C for 30 min. Sections were dehydrated

x SSC/1 mM DTT at 65°C for 30 min. Sections were dehydrated in ascending ethanol, vacuum dried at room temp for 30 min, and then exposed to Dupont Cronex film for several days.

20 Dried slides were then dipped in NTB-2 emulsion (Kodak), and

20 Dried slides were then dipped in NTB-2 emulsion (Kodak), and developed after 6 days.

#### 6.2. RESULTS

#### 6.2.1. GENERATION OF MC4-R DEFICIENT MICE

- The murine MC4-R gene consists of approximately 1 kb of coding sequence contained within a single exon (Figure 1A).

  A targeting vector was designed to delete virtually all MC4-R coding sequence following homologous recombination with the locus in embryonic stem (ES) cells. As shown in Figure 1A,
- 30 the vector consists of a total of approximately 4.5 kb of strain 129/Sv mouse genomic DNA flanking a deletion of 1.5 kb. This deletion extends from the Nco I site located approximately 20 nucleotides downstream of the MC4-R initiation codon to the Hind III site situated approximately
- 35 0.5 kb 3' of the gene. The deleted MC4-R sequences have been replaced by the neo gene under the control of the phosphoglycerate kinase-1 (PGK-1) promoter.

A total of 809 G418-resistant colonies were screened for homologous recombination by Southern blot hybridization of Apa I digested genomic DNA with the flanking probe shown in Figure 3A. One clone showed the predicted 7.6 kb targeted 5 Apa I DNA fragment in addition to the expected 2.2 kb wild type fragment. Injection of this clone into C57BL/6J blastocysts produced several male chimeras which, when bred to C57BL/6J females, transmitted the targeted MC4-R allele to their F1 129/B6 offspring. F1 heterozygotes were interbred 10 and their offspring genotyped by Southern blot hybridization of Apa I or Nco I digested tail DNA with the flanking probe. As described above, Apa I digestion generates a wild type fragment of 2.2 kb and a targeted fragment of 7.6 kb (note that this 7.6 kb is distinct from a background band of 15 slightly lower molecular weight which is present in all samples; Figure 3E). Nco I digestion generates two wild type fragments of 2.7 and 2.9 kb, since the Nco I site is situated within the sequences recognized by the flanking probe (Figure 3). The 2.7 kb Nco I fragment represents genomic 20 sequences extending 3' of the probe which are unaffected by MC4-R targeting, whereas the 2.9 kb band includes the MC4-R gene sequences. Following targeting, this latter fragment is reduced to a 2 kb band diagnostic of the mutated MC4-R allele. As shown in Figure 3E, heterozygous intercrosses 25 produced homozygous mutant, heterozygous and wild type F2 To verify deletion of the MC4-R gene in homozygous mutants, the filters were stripped and rehybridized with an MC4-R probe. No MC4-R hybridization was detected in homozygous mutant mice, whereas the predicted 2.2 kb and 30 5.1 kb Apa I bands (Apa I cuts within the MC4-R gene generating two MC4-R-containing fragments; see Figure 3A) and 2.9 kb Nco I fragment were observed in both heterozygous and

wild type littermates (Figure 3E).

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6.2.2. BODY WEIGHT AND SIZE OF MC4-R DEFICIENT MICE

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weight regulation.

F2 animals were maintained on a chow diet ad libitum and their weights monitored regularly. The weights of MC4-R deficient mice and their wild type littermates were largely 5 indistinguishable for the first 4 weeks of life. However, by approximately 5 weeks of age most of the homozygous mutants, both males and females, were heavier than their wild type siblings of the same sex, and by 7 weeks of age all of the null mutants were heavier than the controls (Figure 4A and 10 4C). By 15 weeks of age, homozygous mutant females were on average twice as heavy as their wild type siblings, while homozygous mutant males were approximately 50% heavier than wild type controls. Weight gain for both males and female mutant mice appeared to be approaching a plateau by about 15 24 weeks of age at which time the weight of female null mice averaged approximately 63 grams (n=3), and males averaged approximately 65 grams (n=8). Mice heterozygous for MC4-R deletion showed a weight gain intermediate to that seen in wild type and homozygous mutant sibs (Figure 4B and 4D), 20 demonstrating a gene dosage effect of MC4-R ablation on body

One of the distinguishing characteristics of the obese yellow phenotype is an increase in skeletal growth. Typically,  $A^{\gamma}$  mice are on average approximately 5% longer than

- 25 their wild type siblings (Castle, 1941, Genetics 26:177191; Carpenter and Mayer, 1958, Am. J. Physiol. 193:499-504). To determine whether mice lacking the MC4-R exhibit enhanced linear growth, body length measurements of F2 progeny were taken at approximately 19 weeks of age (between 132-138).
- 30 days). As shown in Figure 6, MC4-R deficient mice are significantly longer than wild type controls. The mean length of homozygous mutant females is increased approximately 11% relative to wild type F2 mice, and heterozygous females are approximately 7%longer than
- 35 controls. Male homozygotes and heterozygotes are approximately 8% and 2.5% longer than controls, respectively.

#### 6.2.3. FOOD CONSUMPTION

To determine whether food consumption was increased in mice lacking the MC4-R, homozygous mutant females and wild type F2 controls were monitored for food intake over a 2 week 5 period. A' mutants, on a C57BL/6J background, and C57BL/6J controls were also monitored. As previously documented (Frigeri et al., 1988, Endocrinology 113:2097-2105; Shimizu et al., 1989, Life Sciences 45:543-552), A' mice were hyperphagic, eating 36% more than C57BL/6J controls.

10 Similarly, absence of the MC4-R also resulted in a significant increase (46%) in food consumption over wild type F2 controls (Figure 7).

#### 6.2.4. SERUM ANALYSIS

Blood was collected from MC4-R deficient mice and wild type controls over three time intervals (4-8 weeks, 10-14 weeks, 17-23 weeks) and serum assayed for glucose and insulin levels. Serum glucose levels were essentially unchanged in females heterozygous or homozygous for MC4-R 20 deletion, but both heterozygous and homozygous males were hyperglycemic (Figure 8A and 8B). This was first evident for homozygous males at the 10-24 week interval at which time glucose levels were elevated over 2 fold above controls, to 390 mg/dl, but heterozygous mutants showed only a slight 25 elevation of serum glucose at this age. By 17-23 weeks of age, both heterozygous and homozygous male mutant mice showed a doubling of normal serum glucose levels (334 and 361 mg/dl, respectively) relative to controls (156 mg/dl).

Both male and female mutant mice were hyperinsulinemic

30 (Figure 8C and 8D). Nine fold and 5 fold increases in insulin levels were evident in the sera of homozygous mutant females and males, respectively, at 4-8 weeks of age. These levels increased dramatically over time, such that by 17-23 weeks of age the mean concentration of insulin in the serum of homozygous mutant females was approximately 65 ng/ml, and for males approximately 130 ng/ml, representing approximately 60 and 14 fold increases, respectively, over insulin levels

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in F2 wild type controls. Heterozygous mutants were also hyperinsulinemic, although less so than homozygous mutants. For both male and female heterozygotes, a significant difference in insulin levels relative to controls was first observed at the 10-14 week interval; by 17-23 weeks mean insulin levels of heterozygotes were elevated to approximately 10 ng/ml (females) and 85 ng/ml (males).

In addition to glucose and insulin, serum leptin and corticosterone levels were also determined. Leptin levels 10 are elevated in A' mice (Maffei et al., 1995, Nat. Med. 1:1155-1161; Mizuno et al., 1996, Proc. Natl. Acad. Sci. 93:3434-3438), indicating that the syndrome does not result from defects in leptin production. Consistent with postulated role of the hormone in signaling fat depot levels 15 (Campfield et al., 1995, Science 269:546-549; Halaas et al., 1995, Science 269:543-546; Pelleymounter et al., 1995, Science 269:540-543), leptin is also elevated in MC4-R deficient mice (Figure 8E and 8F). At 4-8 weeks of age leptin was elevated 4.5 fold and 1.5 fold in the serum of 20 female and male homozygous mutants, respectively, relative to wild type controls. By 17-23 weeks of age serum leptin levels in females had reached approximately 97 ng/ml, in males approximately 58 ng/ml, representing increases of 6.5 and 2.5 fold, respectively. Heterozygous mice, for the most 25 part, showed leptin levels intermediate between that observed for wild type mice and homozygous mutants.

Since glucocorticoids can profoundly effect weight homeostasis and somatic growth, basal serum corticosterone was measured in three sets of sex-matched littermates, each containing a wild-type, heterozygous, and homozygous mutant animal (Figure 9). No effect of MC4-R gene knockout on basal corticosterone levels was detected.

#### 6.2.5. POMC GENE EXPRESSION

To assess whether the observed effects of MC4-R deletion on weight homeostasis could be attributed to the induction of compensatory changes in the sole known source of ligand for

the MC4-R, the POMC gene, central POMC gene expression was examined in wild type mice, mice heterozygous for MC4-R deletion and homozygous mutants by in situ hybridization (FIG.10A-10F). No new sites of POMC gene expression and no 5 consistent change in the levels of POMC mRNA in its primary site of expression, the arcuate nucleus of the hypothalamus, were detected by this assay. In addition, no gross neuroanatomical defects were observed in thionin-stained brain sections from heterozygous or homozygous mutant MC4-R 10 deficient animals by histological analysis (FIG. 10A-10C).

## 7. EXAMPLE: AGOUTI PROTEIN BINDS DIRECTLY TO MC1-R AND MC4-R

The following example describes experiments

demonstrating that the Agouti protein binds directly to the melanocortin receptors.

#### 7.1. MATERIAL AND METHODS

Human melanocortin receptor 4 (hMC4-r) cDNA, under the control of the CMV promoter, was transfected into the 293 cell line, and stable clones were selected (293/MC4-R). The stable clones were tested for reduction in intracellular cAMP levels in the presence of 5nM agouti protein.

the DEAE-Dextran method. A plasmid containing the Adenovirus VA1 and VA2 RNA genes was used to co-transfect the COS cells to enhance transient protein expression by increasing translational initiation. The MC4-r and the VA1/2 cDNA plasmids were used at a ratio of 10 to 1. Control plates received the VA1/2 plasmid alone. 48 hours post-transfection, the cells were rinsed and culture supernatant containing 15nM of AP-Ag was added. AP-Ag is a truncated agouti protein, containing the cysteine-rich domain tagged with the alkaline phosphatase at its N-terminal. Binding of AP-Ag to transfected cells proceeded at room temperature for 90 minutes with gentle rocking. The cells were then washed 7

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times before fixing and color development using NBT/BCIP substrate.

#### 7.2. RESULTS

When 5 nM agouti protein was added to the 293/MC4-R line, an 18% reduction of intracellular cAMP level, relative to the parental 293 cell line, was observed. The results indicate that MC4-R mediates the agouti-triggered decrease of intracellular cAMP level and confers agouti response to 293 to cells.

N-terminal truncated agouti protein, containing only the cysteine-rich C-terminal domain, retains the antagonizing activity of the full-length agouti. A truncated agouti protein, containing only the cysteine-rich domain and tagged 15 with the alkaline phosphatase at its N-terminal, was used to assay the direct binding between the agouti protein and the MC1 and MC4 receptors in COS7 cells.

The transfection efficiency, as monitored by β-galactosidase reporter plasmid, typically was 12-16%. 10-13% of the MC1-r transfected and 3-4% of the MC4-r transfected COS7 cells bound the agouti protein as determined by AP staining. The observed difference in percentage of positive cell between the MC1-r and MC4-r transfected COS7 cells could be attributed to differences in binding affinity and/or expression level.

To assess the affinity of the agouti protein for the MC1-R receptor, a Scatchard Analysis was performed on MC1-r transfected COS7 cells using culture supernatant containing up to 30 nM AP-Ag. The Kd has been estimated at 20-30nM 30 range.

#### 7.3. DISCUSSION

The dominant agouti alleles that give rise to obesity result from constitutive deregulated synthesis of wild-type 35 agouti protein throughout the animal, and presumably accounts for the other characteristics of the pleiotropic obesity syndrome such as hyperphagia, hyperinsulinemia, and

hyperglycemia. One possible mechanism by which ectopic agouti expression induces obesity is aberrant antagonism of melanocortin receptors, such as MC4-R, expressed in regions of the brain known to be involved in regulating feeding.

The data presented herein demonstrates that the Agouti protein binds directly to MC4-R. Furthermore, as described in Example 6, supra, transgenic mice lacking the MC4-R produce an obesity syndrome that strikingly resembles the agouti syndrome. The recapitulation of many of the features of the agouti syndrome in MC4-R deficient mice demonstrates that antagonism of melanocortin signaling via MC4-R is the primary cause of the agouti obesity syndrome.

# 8. EXAMPLE: IDENTIFICATION AND FUNCTIONAL CHARACTERIZATION OF HUMAN MELANOCORTIN 4 RECEPTOR Ile137Thr MUTANT

Several mutant genes have been identified that cause obesity in mouse; in human, however, only a mutation in β3 adrenergic receptor gene has been inconclusively associated with obesity. The identification of the MC4-R Ile137Thr mutation in obese human subject and the finding of its impaired signaling support MC4-R as a causative genetic factor contributing to human obesity, and validate the receptor as a potential drug target.

#### 8.1. MATERIALS AND METHODS

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8.1.1. <u>DETECTION OF MUTANT MC4-R RECEPTOR</u>
Human genomic DNA isolated from total white blood cells
was amplified by PCR using the following primer pairs:

	•
MC4f1b	5'-TGTAAAACGACGGCCAGTCTGACCCAGGAGGTTAAATC-3'
30 MC4r1b	5'-CAGGAAACAGCTATGACCGCTGCAGATGAAAAAGTACATG-3'

MC412b	5'-TGTAAAACGACGGCCAGTTGCTACGAGCAACTTTTTCTC-3'
MC4r2b	5'-CAGGAAACAGCTATGACCGGTACTGGAGAGCATAGAAG-3'

MC4f3	5'-TGTAAAACGACGGCCAGTTGGTGAGCGTTTCAAATGGAT-3'
MCArs	5'-CAGGAAACAGCTATGACCGAGCCAGCATGGTGAAGAAC-3'

<sup>35</sup> MC4f4 5'-TGTAAAACGACGCCAGTATCTTCTATGCTCTCCAGTAC-3' MC4r4 5'-CAGGAAACAGCTATGACCTTCTGAGGACAAGAGATGTAG-3'

MC4f5b 5'-TGTAAAACGACGGCCAGTTTCTCTCTATGTCCACATGTTC-3'
MC4r5b 5'-CAGGAAACAGCTATGACCGAGTGAAAAAGTCTCTTATGCATG-3'

The following PCR amplification conditions were used:
The 25μL PCR reactions contained a buffer (10 mM Tris-HCl, pH
5 8.3,50mM KCl, 1.5 mM MgCl<sub>2</sub>), the four dNTPs at 100 μM each,
1μM each primer, 0.5U Taq polymerase, and 50-100 ng genomic
DNA. The fragments were amplified in a touchdown PCR (94°C
30 sec, 65°C-55°C 30 sec (-1°C/cycle), 72° 40 sec), followed
by 30 cycles of 94°C 30 sec, 55°C 30 sec, and 72°C 40 sec.

The amplicons were treated with Exonuclease I to remove residual single-stranded primers and Shrimp Alkaline Phosphatase to remove unincorporated dNTPs [Template Purification Kit for Sequencing, Amersham US70995]. The purified fragments were then digested with restriction enzymes MspI and HinfI to yield fragments smaller than 250bp, sizes better suited to analysis by SSCP.

The digested PCR products were diluted 1:9 in loading buffer (95% formamide, 20 mM EDTA, 0.05% bromphenol blue, 0.05% xylene cyanol), heat denatured at 98°C for 2 minutes, 20 and snap-cooled in an ice slurry. 2-3 µL are loaded on a 10% acrylamide gel (50:1 acrylamide:bis-acrylamide) with 10% glycerol and run at 25 watts for 4 hours at 4°C. The gel was stained with SYBR Green I and II to detect both single- and double-stranded DNA fragments, and visualized on a 25 fluorimager.

8.1.2. RECOMBINANT EXPRESSION OF MUTANT MC4-R
Genomic DNA containing the MC4-R wild type and MC4-R
Ile137Thr variant were used as templates for a PCR
30 amplification reaction. Primers flanking the MC4-R coding region were used to PCR the entire MC4 including the MC4-R
Ile137Thr mutation. PCR primers and conditions were as follows:

HMC4 upper primer 5'-CGTAGGATCCATGGTGAACTCCACCCACCTG-3'
35 HMC4 lower primer 5'-AGCCTCGAGTTAATATCTGCTAGACAAGTC-3'
PCR conditions:

96 C for 15 minutes; 94 C for 1 minute; 55 C for 1 minute;

73 C for 2 minutes for 35 cycles; 73 C 10 minutes; 4 C.The resultant 1 kb PCR fragments were subsequently cloned into pCDNA3 (Invitrogen) and sequenced to confirm the correct DNA sequence.

Expression vectors containing human MC4-R wildtype and I137T mutant cDNAs, respectively, were transfected into 293T cells (HEK 293 cells expressing SV40T antigen) using Lipofectamin (BRL). 48 hours later, transfected cells were plated into 96 well plates. After an additional 24 hours, 10 the transfected cells were incubated at 37°C for 15 minutes (or at room temperature for 1 hour) with various malanocortins (BACHEM). Intracellular cAMP levels were then determined using a SPA-based assay (Amersham).

To monitor and normalize the transfection efficiency of the wildtype and mutant cDNAs, an expression vector containing  $\beta$  galactosidase gene were always co-transfected with the MC4-R expression vectors ( $\beta$  galactosidase and MCR-R cDNA at a ratio of 1:9).  $\beta$  galactosidase assay performed 72 hours post-transfection confirmed that the wildtype and 20 I137T mutant MC4-Rs had the same transfection efficiency.

#### 8.2. RESULTS

216 unrelated individuals were screened for sequence variation within the coding region of the MC4R gene. The 216
25 individuals comprised of 96 extreme obese as determined by body mass index (BMI>50), 24 obese (BMI:30-50), 18 intermediates (BMI 25-30), 54 leans (BMI,25) and 24 polycystic ovary syndrome (PCOS) patients. 3 amino acid variants have been found in MC4R in this sample. The three variants are Vallo2Ile (G to A; see FIG. 12A-B), Ilel37Thr (T to C; see FIG. 11A-B) and Thr112Met (C to T; see FIG. 13A-B).

The VallO2Ile substitution was found amongst all categories of BMI within the sample. The following individuals were heterozygous for this variant: 2 extreme

35 obese (BMI=74.2; BMI=57.2), 2 obese (BMI=43; BMI=26), 2 leans (BMI=22.5; BMI+21).

とし 1/039//03909

The Ilel37Thr variant was only found once in an extreme obese person (BMI=57.3). This individual is heterozygous for this variant. The substitution of a threonine (polar) for an isoleucine (non-polar) amino acid may have an effect on the 5 receptor. It is possible that this substitution diminishes the activity of the MC4 receptor contributing to the obesity in this patient.

The Thr112Met variant was only found once (BMI-17.9).

This individual is heterozygous for this variant. The

10 substitution of a methionine (non-polar) for an threonine
(polar) amino acid may have an effect on the receptor. It is
possible that this substitution increases the activity of the
MC4 receptor contribuing to the leanness of this individual.

In order to test whether the Ile137Thr varient was able 15 to function normally, the activity of the Ile137Thr mutant receptor was compared to a wild-type receptor in a signaling assay. Figure 14 compares the response to five endogenous melanocortins,  $\alpha$ -MSH (alpha),  $\beta$ -MSH (beta),  $\gamma$ 1-MS (gamma1),  $\gamma$ 2-MSH (gamma2), and ACTH.

The results demonstrate the impaired signaling of the Ile137Thr mutant receptor as measured by cAMP induction in the presence of various agonists. Compared with the wildtype receptor, the mutant has much lower maximum activation, i.e. lower maximum cAMP level achieved; and it generally has higher EC50, i.e., higher agonist concentration required to reach half maximum activation. The mutant receptor is not

totally inactive in the <u>in vitro</u> assay; but it is marginally active only in the presence of very high agonist concentration that may not be reached under physiological

30 conditions in vivo.

The present invention is not to be limited in scope by the specific embodiments described which are intended as single illustrations of individual aspects of the invention, and functionally equivalent methods and components are within 35 the scope of the invention. Inde d various modifications of the invention, in addition to those shown and described herein will become apparent to those skilled in the art from

the foregoing description and accompanying drawings. Such modifications are intended to fall within the scope of the appended claims.

rc1/U39"//U9909

#### WHAT IS CLAIMED IS:

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1. A method for identifying compounds that regulate body weight, comprising:

- (a) contacting a test compound with a cell which expresses a functional melanocortin 4-receptor, and
- (b) determining whether the test compound activates the melanocortin 4-receptor,

in which test compounds that activate the melanocortin 4receptor are identified as compounds for inducing weight
loss.

- 2. A method for identifying compounds that regulate body weight, comprising:
- (a) contacting a melanocortin peptide in the presence
  and absence of a test compound with a cell which
  expresses a functional melanocortin 4-receptor, and
  - (b) determining whether the test compound inhibits the melanocortin peptide induced activation of the melanocortin 4-receptor,
- 20 in which test compounds that inhibit activation of the melanocortin 4-receptor are identified as compounds for inducing weight gain.
- The method of Claim 1 or 2 in which activation of the
   melanocortin 4-receptor is determined by measuring induction of cAMP.
- The method of Claim 3 in which the cell further contains a reporter gene operatively associated with a cAMP responsive
   element, and induction of cAMP is indicated by expression of the reporter gene.
  - 5. The method of Claim 4 in which the reporter gene is alkaline phosphatase, chloramphenicol acetyltransferase,
- 35 luciferase, glucuronide synthetase, growth hormone, or placental alkaline phosphatase.

- 6. The method of Claim 2 in which the melanocortin peptide is  $\alpha$ -MSH.
- 7. A method for identifying compounds that regulate body 5 weight, comprising:
  - (a) contacting a test compound with a melanocortin 4receptor, and;
  - (b) determining whether the test compound interacts with the melanocortin 4-receptor,
- 10 in which test compounds that interact with the melanocortin 4-receptor are identified as compounds that regulate body weight.
- 8. A method for identifying compounds that regulate body
  15 weight, comprising:
  - (a) contacting a melanocortin peptide in the presence and absence of a test compound with a melanocortin 4-receptor; and
- (b) determining whether the test compound inhibits the interaction of the melanocortin peptide with the melanocortin 4-receptor,

in which test compounds that inhibit the interaction of the melanocortin peptide with the melanocortin 4-receptor are identified as compounds that regulate body weight.

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- 9. The method according the Claim 7 or 8 in which the melanocortin 4-receptor is contained in an isolated membrane or is recombinantly expressed.
- 30 10. The method according to Claim 8 in which the melanocortin peptide is  $\alpha MSH$ .
  - 11. A method for identifying compounds that regulate body weight, comprising:
- 35 (a) contacting a test compound with a cell or cell lysate containing a reporter gene operatively

associated with an melanocortin 4-receptor regulatory lement; and

- (b) detecting expression of the reporter gene product.
- 5 12. A method for identifying compounds that regulate body weight comprising:
  - (a) contacting a test compound with a cell or cell lysate containing melanocortin 4-receptor transcripts; and
- (b) detecting the translation of the melanocortin transcript.
- 13. A method for the treatment of body weight disorders, comprising modulating the activity of the melanocortin 4-15 receptor.
- 14. A method for the treatment of body weight disorders, comprising administering an effective amount of a compound that agonizes or antagonizes the activity of the melanocortin 20 4-receptor.
  - 15. The method of Claim 14 in which the compound activates the melanocortin 4-receptor and induces weight loss.
- 25 16. The method of Claim 15 in which the compound is an agonist or an antibody that binds to and activates the melanocortin 4-receptor.
- 17. The method of Claim 14 in which the compound inhibits 30 activation of the melanocortin 4-receptor and induces weight gain.
- 18. The method of Claim 17 in which the compound is an antibody or a soluble extracellular domain of the35 melanocortin 4-receptor.

- 19. A method for the treatment of body weight disorders, comprising modulating the activity of the melanocortin 4-receptor gene.
- 5 20. A method of the treatment of body weight disorders, comprising administering an effective amount of a compound that decreases expression of the melanocortin 4-receptor and induces weight gain.
- 10 21. The method of Claim 20 in which the compound is an oligonucleotide encoding an antisense or ribozyme molecule that targets melanocortin 4-receptor transcripts and inhibits translation.
- 15 22. The method of Claim 20 in which the compound is an oligonucleotide that forms a triple helix with the promoter of the melanocortin 4-receptor gene and inhibits transcription.
- 20 23. A pharmaceutical formulation for the treatment of body weight disorders, comprising a compound that activates the melanocortin 4-receptor, mixed with a pharmaceutically acceptable carrier.
- 25 24. The pharmaceutical formulation of Claim 23 in which the compound is a melanocortin peptide agonist or an antibody specific for the melanocortin 4-receptor.
- 25. A pharmaceutical formulation for the treatment of body 30 weight disorders, comprising a compound that inhibits the selective activation of the melanocortin 4-receptor, mixed with a pharmaceutically acceptable carrier.
- 26. The pharmaceutical formulation of Claim 25 in which the 35 compound is an antagonist or an antibody specific for the melanocortin 4-receptor.

27. The pharmaceutical formulation of Claim 25 in which the compound is an oligonucleotide encoding an antisense or ribozyme molecule that targets melanocortin 4-receptor transcripts and inhibits translation.

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28. The pharmaceutical formulation of Claim 25 in which the compound is an oligonucleotide that forms a triple helix with the promoter of the melanocortin 4-receptor gene and inhibits transcription.

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- 29. A genetically engineered obese animal in which the melanocortin 4-receptor gene has been inactivated.
- 30. A transgenic animal which expresses a human melanocortin 15 4-receptor gene.
  - 31. A kit comprising a melanocortin 4-receptor or cells expressing the melanocortin 4-receptor packaged in a container.

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- 32. A method for diagnosing body weight disorders, in a mammal comprising measuring MC4-r gene expression in a patient sample.
- 25 33. The method of Claim 32 in which expression is measured by detecting mRNA transcripts of the MC4-r gene.
  - 34. The method of Claim 32 in which expression is measured by detecting the MC4-r gene product.

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- 35. A method for diagnosing body weight disorders in a mammal, comprising detecting a MC4-r gene mutation contained in the genome of the mammal.
- 35 36. The method of Claim 35 in which the mutation is located at nucleotide 137 of the MC4-r gene.

- 37. An isolated DNA consisting of an altered MC4-r sequence having a G at nucleotide 102.
- 38. An isolated DNA consisting of an altered MC4-r sequence 5 having a C at nucleotide 137.
  - 39. An isolated DNA consisting of an altered MC4-r sequence having a T at nucleotide 112.
- 10 40. A nucleic acid probe complementary to human altered MC4-r gene sequences, wherein said nucleic acid probe hybridizes to a mutant MC4-r gene under conditions which prevent hybridizing of said nucleic acid probe to an MC4-r gene having a wild-type sequence.

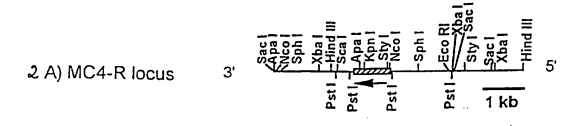
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- 41. The nucleic acid probe of Claim 40 wherein the mutant MC4-r gene has a G at the nucleotide corresponding to base number 102 in the MC4-r gene sequence.
- 20 42. The nucleic acid probe of Claim 40 wherein the mutant MC4-r gene has a C at the nucleotide corresponding to base number 137 in the MC4-r gene sequence.
- 43. The nucleic acid probe of Claim 40 wherein the mutant 25 MC4-r gene has a T at the nucleotide corresponding to base number 112 in the MC4-r gene sequence.
  - 44. A method for identifying compounds that regulate body weight, comprising:
- 30 (a) contacting a test compound with a cell which expresses a mutant melanocortin 4-receptor, and
  - (b) determining whether the test compound activates the melanocortin 4-receptor,

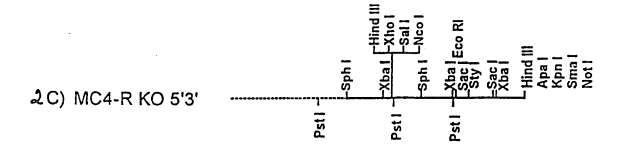
in which test compounds that activate the melanocortin 4-35 receptor are identified as compounds for inducing weight loss.

ECD 1 HSIOKKYLEG DEVEPVSSSS FLRTLLEPOL GSALLTAMNA SCCLPSVOPT HC) нкн HC4 LPNGSEHLOA PFFENOSSSA FCEOVFIKPE IFLSLGIVSL LENILVILAV HC4
HC4
HC2 (ACTH)
HC1 (Q-HSH)
HC1 (Q-HSH)
HC2 (ACTH)
HC3 (Q-HSH)
HC4 LINSTPTAIPQ LQLAANQTGA RCLEVSISIGE LFLSLGLVSL VENALVVATI CD 1 \_II, VANGRILISPH YFFICELAVA DHILVSVSNAL ETIKLATVHS DYLTFEDOFI AXNONLHSPH YFFICSLAVA DHLVSVSNGS ETHILLINS T.DIDAQSFT HC3 FINICILOAPH YFFICSLATS DHLGSLYKIL ENILIILINH GYLKPROSFE HC4 HC1 (Q-HSH) AKHRHLHSPH YCFICCIALS DLLVSGTNVL ETAVILLEA GALVARAAVL CD 2 4 OHHDNIFDSH ICISLVASIC NLLAIAVDRY VTIFYALRYH SIHTVRKALT IIL VNIDAVIDSV ICSSLLASIC SLLSIAVDRY FTIFYALOYH HIHTVKRVGI XC3 TTADDIIDSL FVISLESIF SLSVIAADRY ITIFHALRYH SIVHERETVV HC4 HC1 (Q-HSH) QQLDRVIDVI TCSSHLSSLC FLGATAVDRY ISIFYALRYH SIVTLPRARQ ECD 3 LIVATHYCCG VCGVVFIVYS ESKHVIVCLI THFFAMILIH GTLYVINFLF IISCIMACT VSGILFIIYS DSSAVIICLI THFFTHLALH ASLYYROFTH HC3 VLIVINITET GIGITHVIFS KHVPTVITET SLEPLHLVFI LCLYVRGELL HC4 HC1 (G-HSH) AVANTHVASV VESTLETAYY DHVAVLLCLV VEFLANLVIN AVLYVINGAR ARLHVKRIAA LPPADGVAPO OHSCHKGAVT ITILLGVFIF CHAPFFLHLV CD 3 ARLHIKRIAV LPGTGAI..R OGANEKGAIT LTILIGVFVV CHAPFFLHLI HC3 ARSHTRKIST LPR...... .. ANNEGAIT LTILLGVFIF CHAPFVLHVL HC4 HC1 (G-HSH) ACCHAOGIAR LHKRO.RPVH OGFGLEGAVT LTILLGIFFL CHGPFFLHLT CD 4 350. \_VII \_ ECD 4 LIITCPINPY CICYTAHENT YLVLINCNSV IDPLIYAFRS LELENTFREI FYISCPONPY CVCFMSHFNL YLILINCNSI IDPLTYALRS QELRATFKEI HC3 LHTFCPSHPY, CACYHSLFOV NGHLINGUAV IDPFLYAFRS PELRDAFKKH HC4 KC1 (G-HSH) LIVLOPEHPT CGCIFKNFNL FLALIICHAI IDPLIYAFHS QELRRILKEV LCGCNGHNLG нсэ ICCYPLOGIC DLSSRY HC4 IFCSRYW HC2 (ACTH) HC1 (Q-HSH) LTCSW

F16. 1







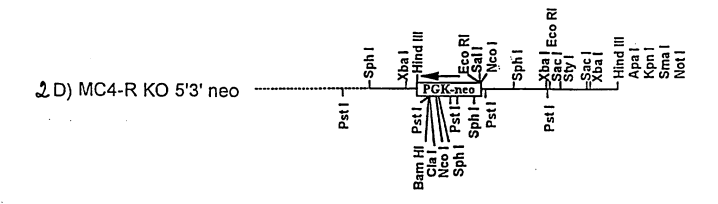
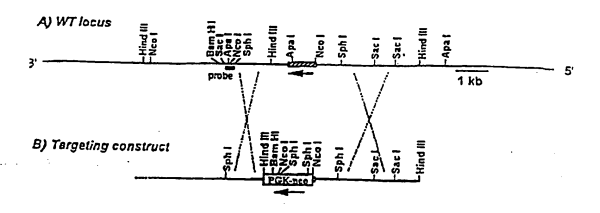
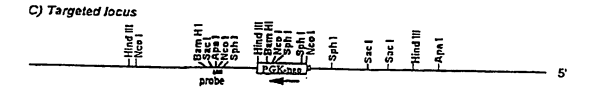


Fig. 2A - 2D





### D) Expected restriction fragment sizes (kb)

Enzyme	M	KQ
Apa I Bam HI Nco I	2.2 -17 2.8 2.6	7.6 2.1 2.8 2.6 1.9
Kpn I Eco RI	4 9.9	1.9 >>4 8.4
CCO 1(1	3.3	0.4

FIGS. 3A-3D

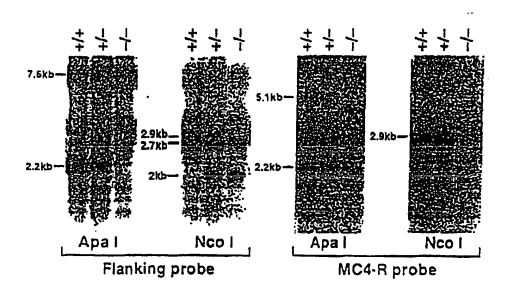
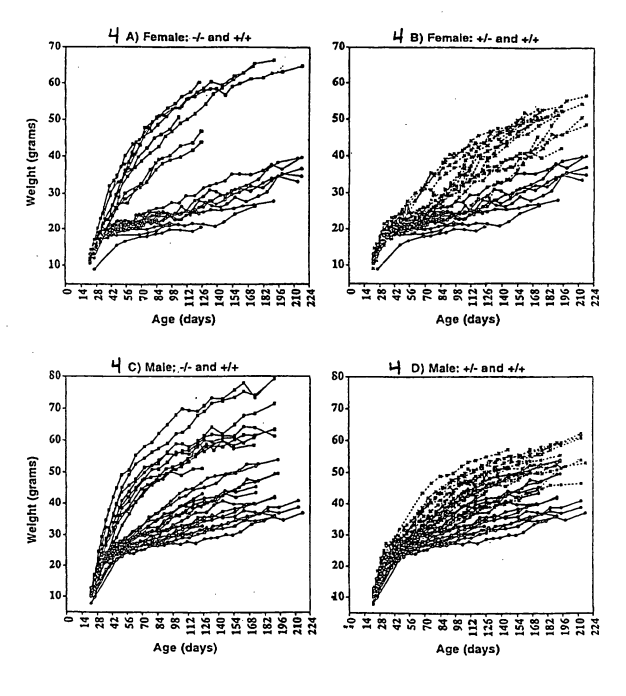
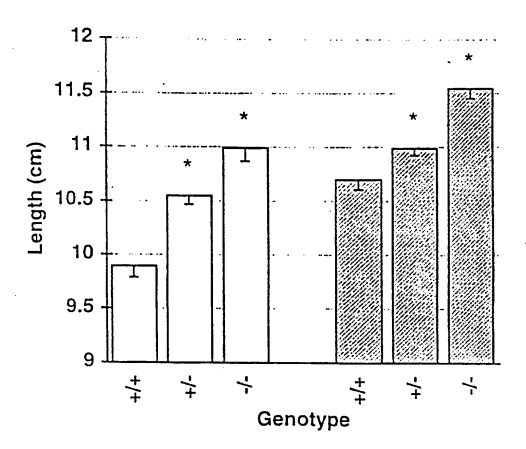


FIG. 3E

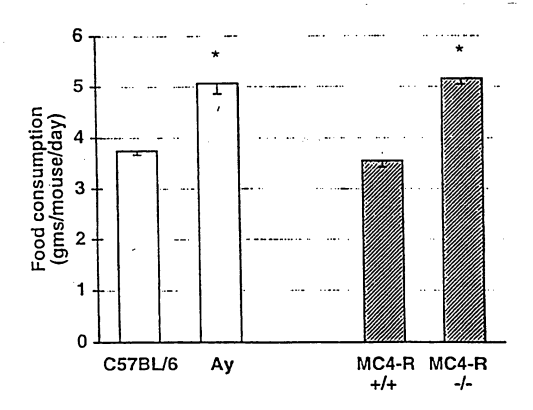


FIGS. 4A-4D

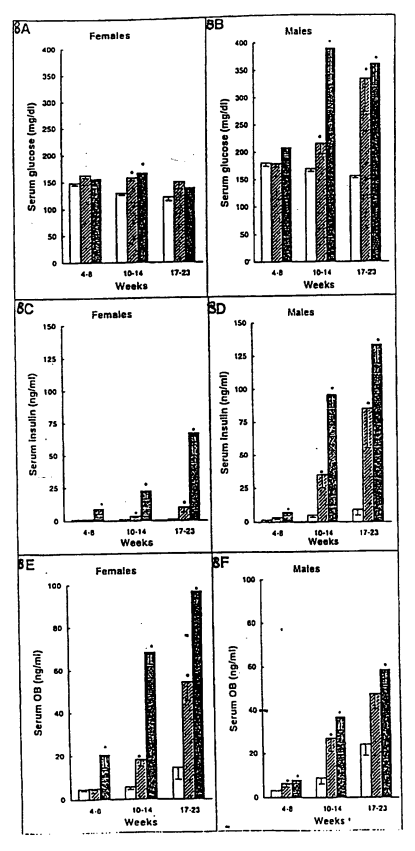
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AGE TTE CGA GAG SEA SEC GAT GTG AGE ATG TGE GGA CAG ATT CCT CTC CCA ATG -110 CCA TGG CAG CTT CAA GGA AAA TTA TTT TGA ACA GAC TTG AAT GCA TAA GAT TAA -286 AGT TAA AGE AGA AGT GAG AAC AAG AAA AAG ACC AGA CTC TTT CAA CTG ACA -217 ATG AAT ATT TTG AAG CCC AAG ATT TTA AAG TGA TGA TGA TTA GAG TGC TAC CTA -178 AAA GAG ACT AAA AAC TCC ATG ACA ACT TTG GA CTT TGA ACA TTT ACT CAC ACC -124 AGG CAT GGC CAT GGC AAT TTT AGC CTC ACA ACT TTC AGA CAG ATT AAG ACT TCG AGG AAA -70 AAG CTG AGA CTC CCT GAC CCC GGA CCC TGG ACA ACT TCA AGC CTC ACA ACT TAA ATC AAT TCA GGG GGA CAC TGG -16
AAT TOT COT GOD AGG ATG GTG AAC TOC ACC CAC CGT GGG ATG CAC ACT TOT GTG 19
HET Val Aun Ser Thr His Arg Gly HET His Thr Ser Leu 13
 CAC CTC TGG-AMC CGC AGC AGT TAC AGA CTG CAC AGC AAT GCC AGT GAG TCC CTT 91
His Lou TEP Ask Arg Ser Ser Tyr Arg Lou His Ser Ask Ala Ser Glu Ser Leu 11
 GGA ANA GGC TAC TOT GAT GGA GGG TGC TAC GAG CAA CTT TIT GTC TCT CCT GAG 147
Gly Lys Gly Tyr Ser Asp Gly Gly Cys Tyr Glu Glu Leu Phe Val Ser Pro Glu 49
GTG TIT GTG ACT CTG GGT GTC ATC AGC ITG TTG GAG AAT ATC TIA GTG ATT GTG 201 Val Phe Val Thr Leu Gly Val Ile Ser Leu Leu Glu Asn Ile Leu Val Ile Val 67
 GCA ATA GCC ANG ANC ANG ANT CTG CAT TCA CCC ATG TAC TTT TTC ATC TGC'AGC 255 Ala Ile Ala Lys Asn Lys Asn Leu His Ser Pro MET Tyr Phe Phe Ile Cys Ser 85
 TTG GCT GTG GCT GAT ATG CTG GTG AGC GTT TCA AAT GGA TCA GAA ACC ATT ATC 309
Leu Ale Val Ale AED MCT Leu Val Ser Val Ser Asn Gly Ser Glu Thr Ile 11e 101
 ATC ACC CTA TTA AAC AGT ACA GAT ACG GAT GCA CAG AGT TTC ACA GTG AAT ATT 361
The The Leu Leu Ash Ser The Asp The Asp Ala Gin Ser Phe The Val Ash Ile 171
 GAT AAT GTC ATT GAC TCG GTG ATC TGT AGC TCC TTG CTT GCA TCC ATT TGC AGC 417 ASP ASP Val Ile Asp Ser val Ile Cys Ser Ser Leu Leu Ale Ser Ile Cys Ser 139
 CTG CTT TCA ATT SCA GTG GAC AGG TAC TTT ACT ATC TTC TAT GCT CTC CAG TAC 471
Leu Leu Ser Ile Ale Val Asp Arg Tyr Phe Thr Ile Phe Tyr Ale Leu Gln Tyr 157
  CAT AAC ATT ATG ACA GIT AAG CGG GIT GGG ATC AGC ATA AGT TGT ATC TGG GCA 525
Ris Asn Ile HET Thr Val Lys Arg Val Gly Ile Ser Ile Ser Cys Ile Trp Ala 175
  GCT TGC ACG GTT TCA GGC ATT TTG TTC ATC ATT TAC TCA GAT AGT AGT GCT GTC 579
Ale Cys The Val Ser Gly Tlo Law Pho Ile Ile Tyr Ser Asp Ser Ser Ale Val 191
  ATC ATC TGC CTC ATC ACC ATG TTC TTC ACC ATG CTG GCT CTC ATG GCT TCT CTC 633 Ile Ile Cys Leu Ile Thr MCT Phe Phe Thr MCT Leu Als Leu MCT Als Ser Leu 211
  TAT GTC CAC ATG TTC GTG ATG GCC AGG CTT CAC ATT AAG AGG ATT GCT GTC CTC 687
TYP Val His HOT Phe Leu HOT Ale Arg Leu His Ile'Lys Arg Ile Ale Val Leu 229
  CCC GGC ACT GGT GCC ATC CGC CAA GGT GCC AAT ATG AAG GGA GCG ATT ACC TTG 741
Pro Gly The Gly Ala 11s Arg Gln Gly Ala Asn MET Lys Gly Ala 11s The Lau 247
  ACC ATC CTG ATT GGC GTC TTT GTT GTC TGC TGG GCC CCA TTC TTC CTC CAC TTA 795
Thr Ile Leu Ile Gly Val Phe Val Val Cys Trp Ala Pro Phe Phe Leu His Leu 265
  ATA TTC TAC ATC TCT TGT CCT CAG AAT CCA TAT TGT GTG TGC TTC ATG TCT CAC 8(9) Ile Phe Tyr Ile Ser Cys Pro Gin Asn Pro Tyr Cys Val Cys Phe MET Ser Mis 281
   VII
TIT AAC TIG TAT CTC ATA CTG ATC ATG TGT AAT TCA ATC ATC GAT CCT CTG ATT 903
Pho Asn Leu Tyr Lou I's Leu I's MET Cys Asn Ser I's Asp Pro Lou I's 301
   TAT OCA CTC COG AGT CAA GAA CTG AGG AAA ACT TTC AAA GAG ATC ATC TGT TOC 957
TYR Als Lou Arg Ser Gln Glu Leu Arg LyseThr Phe Lys Glu Ile Ile Cys Cys 319
   TAT CCC CTG GGA GGC CTT TGT GAC TTG TCT AGC AGA TAT TAA ATG GGG ACA GAG 996
Tyr Pro Leu Gly Gly Leu Cys Amp Leu Sor Ser Arg Tyr 1322
   CAE GCA ATA TAG GAA CAT GCA TAA GAG ACT TIT TCA CTC TTA CCC TAC CTG AAT 1050 ATT GTA CTT CTG CAA CAG CTT TCT-CCT TGT ACG GTA CTG GTT GAG ATA TCC 1108 ATT GTG TAA ATT TAA GCC TAT GAT TTT TAA TGA GAA AAA ATG CCC ACT CTC TGT 1162 ATT ATT TCC AAT GTC ATG CTA CTT TTT TGC CCA TAA AAT ATG AAT CTA TGT TAT 1216 AGG TTG TAG GCA CTG TGG ATT TAC AAA AAG AAA ACT CCT TAT TAA AAG CTT
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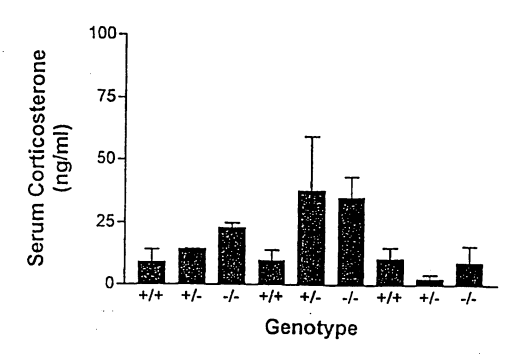
F16.6



F16. 7



FIGS 8A-8F 9/18



F16. 9

D E ÂRC ÂRC

FIGS. 10A - 10F

11/18

	gtg Val															48
aac Asn	Arg Cgc	agc Ser	agt Ser 20	tac Tyr	aga Arg	ctg Leu	cac His	agc Ser 25	aat Asn	gcc Ala	agt Ser	gag Glu	ser 30	ctt Leu	gga Gly	96
aaa Lys	Gly	tac Tyr 35	tct Ser	gat Asp	gga Gly	Gly ggg	tgc Cys 40	tac Tyr	gag Glu	caa Gln	ctt Leu	ttt Phe 45	gtc Val	tct Ser	cct Pro	144
gag Glu	gtg Val 50	ttt Phe	gtg Val	act Thr	ctg Leu	ggt Gly 55	gtc Val	atc Ile	agc Ser	ttg Leu	ttg Leu 60	gag Glu	aat Asn	atc Ile	tta Leu	192
gtg Val 65	att Ile	gtg Val	gca Ala	ata Ile	gcc Ala 70	aag Lys	aac Asn	rys Fys	aat Asn	ctg Leu 75	cat His	tca Ser	ccc Pro	atg Met	tac Tyr 80	240
ttt Phe	ttc Phe	atc Ile	tgc Cys	agc Ser 85	ttg Leu	gct Ala	gtg Val	gct Ala	gat Asp 90	atg Met	ctg Leu	gtg Val	agc Ser	gtt Val 95	tca Ser	288
aat Asn	Gly	tca Ser	gaa Glu 100	acc Thr	gtt Val	atc Ile	atc Ile	acc Thr 105	cta Leu	tta Leu	aac Asn	agt Ser	aca Thr 110	gat Asp	acg Thr	336
gat Asp	gca Ala	cag Gln 115	agt Ser	ttc Phe	aca Thr	gtg Val	aat Asn 120	att Ile	gat Asp	aat Asn	gtc Val	att Ile 125	yab	tcg Ser	gtg Val	384
atc Ile	tgt Cys 130	agc Ser	tcc \$er	ttg Leu	ctt Leu	gca Ala 135	tcc Ser	act Tbr	tgc Cys	agc Ser	ctg Leu 140	ctt Leu	tca Ser	Ile	gca Ala	432
gtg Val 145	gac Asp	agg Arg	tac Tyr	ttt Phe	act Thr 150	atc Ile	ttc Phe	tat Tyr	gct Ala	ctc Leu 155	cag Gln	tac Tyr	cat Kis	aac Asn	att Ile 160	480
atg Met	aca Thr	gtt Val	aag Lys	cgg Arg 165	gtt Val	GJA aaa	atc Ile	agc Sex	ata Ile 170	agt Ser	tgt Cys	atc Ile	tgg Trp	gca Ala 175	gct Ala	528
tgc Cys	acg Thr	gtt Val	tca Ser 180	ggc	att Ile	ttg Leu	ttc Phe	atc Ile 185	att Ile	tac Tyr	tca Ser	gat Asp	agt Ser 190	agt Ser	gct Ala	576
gtc	atc	atc	tgc	ctc	atc	acc	atg	ttc	ttc	acc	atg	ctg	gct	ctc	atg	624

FIG. 11 A 12/18

Val	Ile	Ile 195	Cys	Leu	Ile	Thr	Met 200	Phe	Phe	Thr	Met	Leu 205	Ala	Leu	Met	
						atg Met 215									aag Lys	672
	Ile					GJÀ ââc									aat Aen 240	720
						ttg Leu									gtc Val	768
						ctc Leu									cct Pro	81€
						tge Cys									ctc Leu	864
ata Ile	ctg Leu 290	atc Ile	atg Met	tgt Cys	aat Asn	tca Ser 295	atc Ile	atc Ile	gat Asp	cct Pro	ctg Leu 300	att Ile	tat Tyr	gca Ala	ctc Leu	912
						aaa Lys									tat Tyr 320	960
	_				_	gac Asp	_		-	-			•			999

FIG. 11B

Met	y gro	g aac 1 Asr	Ser	Thr	His	: cgt : Arg	. GJA	atg Met	His 10	Thr	: tct :Ser	Leu	y Cad His	c cto s Lev	tgg Trp	4
aac Asn	cgo Arg	ago g Ser	e agt Ser 20	Tyr	aga Arg	ctg Leu	cac His	ago Ser 25	Asn	gcc	agt Ser	gag	s tcc Ser 30	Lev	gga Gly	9 (
aaa Lys	ggc Gly	tac Tyr 35	Ser	gat Asp	gga Gly	Gly	tgc Cys 40	tac	gag Glu	caa Gln	ctt Leu	ttt Phe 45	Val	tct Ser	cct Pro	144
ga <b>g</b> Glu	gtg Val 50	Pne	gtg Val	act Thr	ctg Leu	ggt Gly 55	gtc Val	ato Ile	agc Ser	ttg Leu	ttg Leu 60	gag Glu	aat Asn	atc Ile	tta Leu	192
gtg Val 65	att	gtg Val	gca Ala	ata Ile	gcc Ala 70	aag Lys	aac Asn	aag Lys	aat Asn	ctg Leu 75	cat	tca Ser	ccc	atg Met	tac Tyr 80	240
ttt	ttc Phe	atc Ile	tgc Cys	agc Ser 85	ttg Leu	gct Ala	gtg Val	gct Ala	gat Asp 90	atg Met	ctg Leu	gtg Val	agc Ser	gtt Val 95	tca Ser	288
aat Asn	Gly	tca Ser	gaa Glu 100	acc Thr	att Ile	atc Ile	atc Ile	acc Thr 105	cta Leu	tta Leu	aac Asn	agt Ser	aca Thr 110	gat Asp	acg Thr	336
gat Asp	gca Ala	cag Gln 115	agt Ser	ttc Phe	aca Thr	gtg Val	aat Asn 120	att Ile	gat Asp	aat Asn	gtc Val	att Ile 125	gac Asp	tcg Ser	gtg Val	384
atc Ile	tgt Cys 130	agc Ser	tcc Ser	ttg Leu	ctt Leu	gca Ala 135	tcc Ser	att Ile	tgc Cys	agc Ser	ctg Leu 140	ctt Leu	tca Ser	att Ile	gca Ala	432
gtg Val 145	gac Asp	agg Arg	tac Tyr	ttt Phe	act Thr 150	atc Ile	rtc Phe	tat Tyr	gct Ala	ctc Leu 155	cag Gln	tac Tyr	cat His	aac Asn	att Ile 160	480
atg Met	aca Thr	gtt Väl	aag Lys	cgg Arg 165	gtt Val	GJA aaa	atc Ile	agc Ser	ata Ile 170	agt Ser	tgt Cys	atc Ile	tgg Trp	gca Ala 175	gct Ala	528
tgc Cys	acg Thr	gtt Val	tca Ser 180	GJÅ ââc	att Ile	ttg Leu	Phe	atc Ile 185	att Ile	tac Tyr	tca Ser	Asp	agt Ser 190	agt Ser	gct Ala	576
gtc	atc	atc	tgc	ctc	atc	acc	atg	ttc	ttc	acc	atg (	ctg	gct	ctc	atg	624

FIG. 12A 14/18

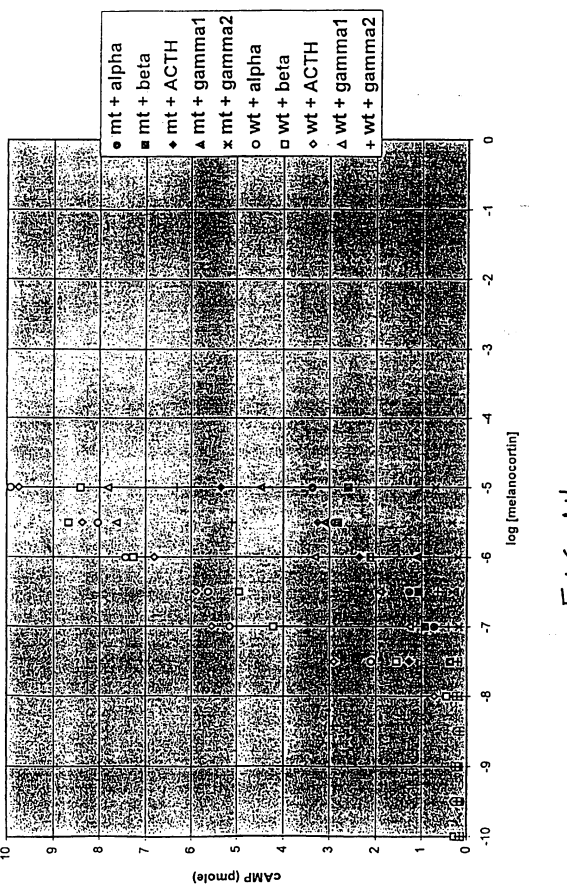
Val Ile Ile Cys Leu Ile Thr Met Phe Phe Thr Met Leu Ala Leu Met get tet etc tat gte cae atg tte etg atg gee agg ett cae att aag 672 Ala Ser Leu Tyr Val His Met Phe Leu Met Ala Arg Leu His Ile Lys 220 agg att gct gtc ctc ccc ggc act ggt gcc atc cgc caa ggt gcc aat 720 Arg Ile Ala Val Leu Pro Gly Thr Gly Ala Ile Arg Gln Gly Ala Asn 235 atg aag gga gcg att acc ttg acc atc ctg att ggc gtc ttt gtt gtc Met Lys Gly Ala Ile Thr Leu Thr Ile Leu Ile Gly Val Phe Val Val 250 tgc tgg gcc cca ttc ttc ctc cac tta ata ttc tac atc tct tgt cct 816 Cys Trp Ala Pro Phe Phe Leu His Leu Ile Phe Tyr Ile Ser Cys Pro cag aat cca tat tgt gtg tgc ttc atg tct cac ttt aac ttg tat ctc Gln Asn Pro Tyr Cys Val Cys Phe Met Ser His Phe Asn Leu Tyr Leu ata ctg atc atg tgt aat tca atc atc gat cct ctg att tat gca ctc 912 Ile Leu Ile Met Cys Asn Ser Ile Ile Asp Pro Leu Ile Tyr Ala Leu 290 cgg agt caa gaa ctg agg aaa acc ttc aaa gag atc atc tgt tgc tat 960 Arg Ser Gln Glu Leu Arg Lys Thr Phe Lys Glu Ile Ile Cys Cys Tyr 315 ccc ctg gga ggc ctt tgt gac ttg tct agc aga tat taa 999 Pro Leu Gly Gly Leu Cys Asp Leu Ser Ser Arg Tyr Stop 325

FIG. 12B

	y y cy t Val															4 8
	cgc Arg															96
	ggc Gly															144
	g gtg Val 50											Glu				192
gtg Val 65	att. Ile	gtg Val	gca Ala	ata Ile	gcc Ala 70	aag Lys	aac Asn	aag Lys	aat Asn	ctg Leu 75	cat His	tca Ser	ccc Pro	atg Met	tac Tyr 80	240
Phe	ttc Phe	atc Ile	tgc Cys	agc Ser 85	ttg Leu	gct Ala	gtg Val	gct Ala	gat Asp 90	atg Met	ctg Leu	gtg Val	agc Ser	gtt Val 95	tca Ser	288
	gga Gly															336
gat Asp	gca Ala	cag Gln 115	agt Ser	ttc Phe	aca Thr	gtg Val	aat Asn 120	att Ile	gat Asp	aat Asn	gtc Val	att Ile 125	gac Asp	tcg Ser	gtg Val	384
atc Ile	tgt Cys 130	agc Ser	tcc Ser	ttg Leu	ctt Leu	gca Ala 135	tcc Ser	att Ile	tgc Cys	agc Ser	ctg Leu 140	ctt Leu	tca Ser	att Ile	gca Ala	432
gtg Val 145	gac	agg Arg	tac Tyr	ttt Phe	act Thr 150	atc Ile	ttc Phe	tat Tyr	gct Ala	ctc Leu 155	cag Gln	tac Tyr	cat His	aac Asn	att Ile 160	480
atg Met	aca Thr	gtt Val	aag Lys	cgg Arg 165	gtt Val	Gly 999	atc Ile	agc Ser	ata Ile 170	agt Ser	tgt Cys	atc Ile	tgg Trp	gca Ala 175	gct Ala	528
cye Cye	acg Thr	gtt Val	tca Ser 180	G]Å aac	att Ile	ttg Leu	ttc Phe	atc Ile 185	att Ile	tac Tyr	tca Ser	gat Asp	agt Ser 190	agt Ser	gct Ala	576
atc	250	atc	tac	ctc	arc	200	ato	ttc	ttc	366	3.50					

Va1	Ile	Ile 195	Cys	Leu	lle	Thr	Met 200	Phe	Phe	Thr	Met	Leu 205	Ala	Leu	Met	
Ala	tct Ser 210	ctc Leu	tat Tyr	gtc Val	His	atg Met 215	ttc Phe	ctg Leu	atg Met	gcc Ala	agg Arg 220	ctt Leu	cac His	att Ile	aag Lys	672
agg Arg 225	att Ile	gct Ala	gtc Val	Leu	ccc Pro 230	Gly Ggc	act Thr	ggt Gly	gcc Ala	atc Ile 235	cgc Arg	caa Gln	ggt Gly	gcc Ala	aat Asn 240	720
atg Met	aag Lys	gg <b>a</b> Gly	gcg	att Ile 245	acc Thr	ttg Leu	acc Thr	atc Ile	ctg Leu 250	att Ile	GJÀ āãc	gtc Val	ttt Phe	gtt Val 255	gtc Val	768
tgc Cys	Trp Trgg	gcc Ala	cca Pro 260	Phe	ttc Phe	ctc Leu	cac His	tta Leu 265	ata Ile	ttc Phe	tac Tyr	atc Ile	ser 270	- 3 -	cct Pro	816
cag Gln	aat Asn	cca Pro 275	Tyr	tgt Cys	gtg Val	cya	ttc Phe 280	met	tct Ser	cac His	ttt Phe	aac Asn 285		tat Tyr	Leu	864
ata Ile	ctg Leu 290	Ile	atg Met	tgt Cys	aat Asn	tca Ser 295	TIG	atc Ile	gat Asp	Pro	t ctg Lev 300		tat Tyr	gca Ala	ctc Leu	912
cgg Arg 305	Ser	caa Glr	ı gaa	a ctg ı Lev	agg Arg 310	LLYS	acc Thr	tto Phe	: aaa : Lys	gag Glu 319		ato	tgi Cy:	tgo S Cys	tat Tyr 320	960
ccc	cto Lev	r GJ:	y Gl	c ctt y Lei 329	7 CA	s Asi	tto Lev	ı Se:	t ago r Sea 330		a ta g Ty:	t ta r St	a op			999

F16. 13B



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#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/09969

A. CLA	SSIFICATION OF SUBJECT MATTER		·
	:Please See Extra Sheet.		
	:Please See Extra Sheet. o International Patent Classification (IPC) or to both	national classification and IPC	
	DS SEARCHED		
Minimum d	ocumentation searched (classification system followed	by classification symbols)	
U.S. :	424/143.1, 145.1; 435/6, 7.1, 8, 346; 514/2, 44; 530	0/312, 399; 536/23.5, 24.5	
Documentat	ion searched other than minimum documentation to the	extent that such documents are included	in the fields searched
Electronic d	ata base consulted during the international search (na	me of data base and, where practicable	, search terms used)
APS; STI	N medline, embase, biosis, scisearch melanocortin, receptor		
C. DOC	UMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where ar	propriate, of the relevant passages	Relevant to claim No.
×	LU, D. et al. Agouti protein		2-3, 6-10
Υ .	melanocyte-stimulating-hormone October 1994, Vol. 371, page document.		1, 4-5, 11, 13- 14, 17, 19, 23, 25, 31
Y	ALBANESE, C. et al. Developme using a recombinant human FSI responsive luciferase reporter gene February 1994, Vol. 101, pag document.	H receptor and a cAMP e. Molec. Cell. Endocrinol. es 211-219, see entire	4-5, 11, 31
X Furth	ner documents are listed in the continuation of Box C	See patent family annex.	·.
*A* dox	ecial categories of cited documents:  cument defining the general state of the art which is not considered be of particular relevance	"I" later document published after the inte date and not in conflict with the applic principle or theory underlying the inv	ation but cited to understand the ention
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*O* dox	ecial reason (as specified)  current referring to an oral disclosure, use, exhibition or other  ans	considered to involve an inventive combined with one or more other suc being obvious to a person skilled in the	step when the document is h documents, such combination
	cument published prior to the international filing date but later than priority date claimed	*&* document member of the same patent	family
	actual completion of the international search	Date of mailing of the international sea	arch report
06 AUGU	JST 1997	1 0 SEP 1997	
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Facsimile N	o. (703) 305-3230	Telephone No. (703) 308-0196	
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International application No. PCT/US97/09969

C (Continue	tion). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relev	ant passages	Relevant to claim No
Y	ALVARO, J.D. Cloning of the rat melanocortin 4 rec brain and its regulation by opiates. Diss. Abstracts Int 1996. Vol. 56, No. 8, page 4165-B, especially DA954 entire abstract.	. February	12
ľ	MAURI, A. et al. Melanocortins and opioids modulate postnatal growth in rats. Regulatory Peptides. July 1959, pages 59-66, see entire document.	e early 195, Vol.	12
Y	ADAN, R.A.H. et al. Identification of antagonists for melanocortin MC3, MC4 and MC5 receptors. Eur. J. 1994, Vol. 269, pages 331-337, see entire document.	Pharmacol.	1-12
. <b>,</b> P	KESTERSON, R.A. et al. Induction of neuropeptide expression in the dorsal medial hypothalamic nucleus ir models of the agouti obesity syndrome. Molec. Endoci 1997, Vol. 11, No. 5, pages 630-637, see entire docum	n two rinol. May	1-44
	HUSZAR, D. et al. Targeted disruption of the melano receptor expression results in obesity in mice. Cell. 1 1997, Vol. 88, No. 1, pages 131-141, see entire documents	O January	1-44
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## INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/09969

. CLASSIFICATION OF SUBJECT MATTER:

A61K 38/16, 39/395, 48/00; C07H 21/04; C12N 15/11; C12Q 1/68; G01N 33/53; C12Q 1/25, 1/66, 1/68

\. CLASSIFICATION OF SUBJECT MATTER:

424/143.1, 145.1; 435/6, 7.1, 8, 346; 514/2, 44; 530/312, 399; 536/23.5, 24.5